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1988 HANFORD RIVERBANK SPRINGS CHARACTERIZATION REPORT

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SUMMARY

This report presents the results of a special study undertaken to characterize the riverbank springs (i.e., ground-water seepage) entering the Columbia River along the Hanford Site. Radiological and nonradiological analyses were performed. River water samples were also analyzed from upstream and downstream of the Site as well as from the immediate vicinity of the springs. In addition, irrigation return water and spring water entering the river along the shoreline opposite Hanford were analyzed.

Hanford-origin contaminants were detected in spring water entering the Columbia River along the Hanford Site. The type and concentrations of contaminants in the spring water were similar to those known to exist in the ground water near the river. The location and extent of the contaminated discharges compared favorably with recent ground-water reports and predictions. Spring discharge volumes remain very small relative to the flow of the Columbia. Downstream river sampling demonstrates the impact of ground-water discharges to be minimal, and negligible in most cases.

Radionuclide concentrations were below U.S. Department of Energy Derived Concentration Guides (DCGs) with the exception of ⁹⁰Sr near the 100-N Area. Tritium, while below the DCG, was detected at concentrations above the U.S. Environmental Protection Agency drinking water standards in several springs. All other radionuclide concentrations were below drinking water standards. Nonradiological contaminants were generally undetectable in the spring water. River water contaminant concentrations, outside of the immediate discharge zones, were below drinking water standards in all cases.

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INTRODUCTION

This report presents the results of a special study performed by the Surface Environmental Monitoring Project (SEMP) to investigate the radiological and nonradiological characteristics of the riverbank springs entering the Columbia River along the Hanford Site shoreline. The SEMP is conducted by Pacific Northwest Laboratory (PNL) for the U.S. Department of Energy (DOE).

Routine SEMP monitoring activities provide a historical record of radionuclide concentrations and radiation levels attributable to natural causes, worldwide fallout, and Hanford Operations. Data are also collected to monitor levels of nonradiological contaminants at the Hanford Site and in the Columbia River. In addition to routine monitoring activities, special studies are also conducted periodically to enhance the understanding of specific aspects of the Hanford environment. The special study described herein was performed during 1988 to supplement the routine monitoring program and to follow up on previously conducted studies investigating similar environmental conditions.

BACKGROUND INFORMATION

The Hanford Site is located in southeastern Washington State, occupying an area of approximately 560 square miles. The Site lies approximately 170 miles southeast of Seattle, Washington, 125 miles southwest of Spokane, Washington, and 200 miles northeast of Portland, Oregon (Figure 1). The Columbia River flows through the northern portion of the Hanford Site and forms part of the Site's eastern boundary. The Rattlesnake Hills, with elevations in excess of 3200 feet, form part of the southern boundary of the Site. The Columbia River Basalt Group, the Ringold Formation, and a series of glaciofluvial sands and gravels known informally as the Hanford sediments are the predominant geological units. Both confined and unconfined aquifers exist beneath the Hanford Site.

The Hanford Site was originally established in 1943 for the production of plutonium for use in nuclear weapons. As a result of operations at

Hanford, large volumes of wastewater were generated and discharged to the ground. The disposal of this liquid effluent to the ground has had a considerable impact on the ground water in the unconfined aquifer beneath the Site. Water levels have been influenced and the local flow patterns, generally from the recharge areas in the west to the discharge areas (primarily the Columbia River) in the east, have been altered. Because of the continual variation in both the volumes and make-up of the wastewater, the movement of the ground water and its associated contaminants have changed with time. In addition, the discharge locations of the contaminated ground water into the Columbia River have expanded over time, encompassing a larger segment of the shoreline.

The Ground-Water Protection and Monitoring Project (GWPMP), formerly the Hanford Ground-Water Surveillance Program, is responsible for monitoring the ground water beneath the Hanford Site. Monitoring is performed via a network of sampling wells located throughout the Site. Sample results are reported in a series of semi-annual and annual ground-water monitoring reports. Recently, the ground-water monitoring and surface environmental monitoring data have been combined in a single Hanford Site environmental monitoring report (Price 1986). While this program has been primarily interested historically in radioactive pollutants in the ground water, nonradiological contaminants have also been monitored during the past few years. In addition to the routine sitewide ground-water monitoring project, several hazardous waste ground-water monitoring compliance projects are ongoing at various locations on the Site that provide information relative to contaminant concentrations in the ground water beneath the Site. Two such projects are presently being conducted in areas near the river, one in the 100-H Area and the other in the 300 Area. These projects provide extensive information about the contaminants in the ground water near the river along these areas (Schalla et al. 1988; Liikala et al. 1988).

Monitoring data have shown several contaminants to be present in the ground water beneath waste disposal sites. The data also indicate that several of these contaminants are mobile in the ground-water system and travel at various rates through the unconfined aquifer, eventually to discharge to the Columbia River. Estimates of ground-water contaminant travel times, made

since the early 1950s, were recently summarized by Freshley and Graham (1988). Tritium and nitrate are the primary constituents used in determining the extent of the contaminated ground water on the Hanford Site since they are present in easily measurable quantities and they move through the ground water virtually unimpeded. Figures 2 and 3 show tritium and nitrate concentrations in the unconfined aquifer during 1988, illustrating the migration of contaminants away from waste disposal areas. The primary areas where contaminated ground water is discharging into the Columbia River are also evident in these figures.

The SEMP is responsible for the monitoring of the surface water on and around the Hanford Site. Columbia River monitoring has been performed at Hanford since 1945, shortly after the startup of the original plutonium production reactors. Samples have been collected routinely from several locations over the years, with the primary emphasis of the program focused on the evaluation of the potential dose to those persons using and/or consuming the river water. In addition, special studies have investigated the mixing characteristics of the river and the dispersion of contaminants entering the river along the Hanford Reach. The springs, or ground-water seepage, are also sampled periodically as part of the SEMP.

The seepage of ground water into the Columbia River has been known to exist for many years. Spring discharges were documented along the Hanford Reach long before the startup of Hanford operations (Jenkins 1922). These relatively small springs flow intermittently, apparently influenced primarily by the changes in the river level. During periods of high river stage, the flow of ground water may be temporarily reversed with river water infiltrating the riverbank (Raymond and Brown 1963). This phenomena, referred to as bank storage, is a key factor in sample collection and in data interpretation. The interface between the ground water and the Columbia River is highly complex and not well defined. Seepage above the river level is considered to be just a fraction of the total amount of ground water entering the river along the Hanford Reach. The exchange of ground-water contaminants at depth between the ground water and the river through submerged seepage is not well understood.

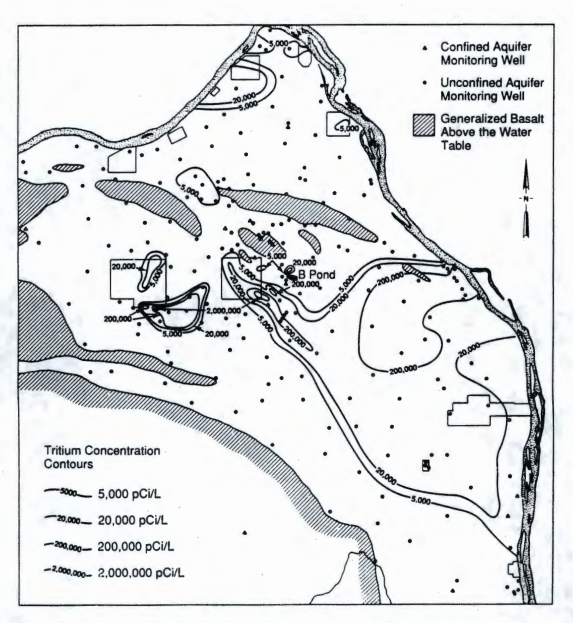


FIGURE 2. Trilium (3H) Concentrations in the Hanford Site Unconfined Aquifer in 1988 (Jaquish and Bryce 1989)

The volume of the ground-water discharge to the river along the entire Hanford Reach has not been quantified. However, estimates of the ground-water discharge in specific areas along the Site have been reported. The N-Springs, adjacent to the 100-N Area, discharged approximately

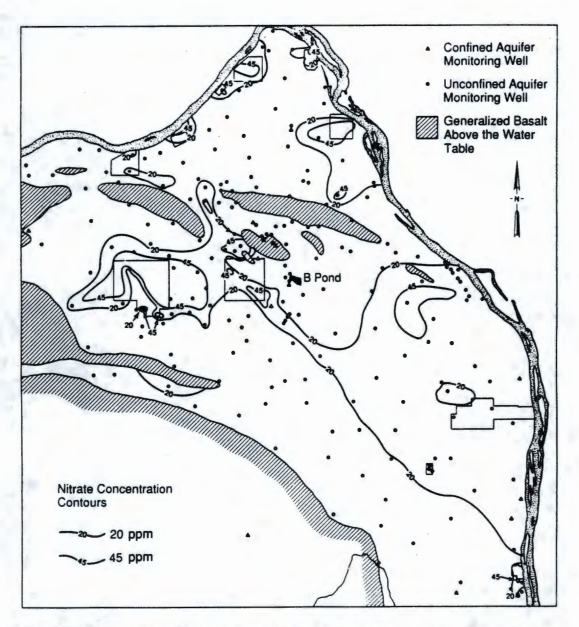


FIGURE 3. Nitrate (NO₃) Concentrations in the Hanford Site Unconfined Aquifer in 1988 (Jaquish and Bryce 1989)

14,700,000 $\rm ft^3$ during 1987, or an average flowrate over the year of approximately 0.5 $\rm ft^3/s$ (Rokkan 1988). The contaminated ground-water discharge to the river near the Hanford Townsite, as a result of past waste disposal practices in the 200 Areas, has been estimated to be approximately 3.0 $\rm ft^3/s$

(Cline et al. 1985). These two areas, 100-N Area and the Hanford Townsite, have been identified as major discharge zones for contaminated Hanford ground water. Based on these estimates, it is apparent that the total flow of ground water into the Columbia River is very small when compared to the flow of the Columbia River. Recent annual average river flow rates have ranged from 100,000 to 120,000 ft 3 /s. The long-term average annual flow at Priest Rapids Dam, based on 60 years of record, is 120,000 ft 3 /s (McGavock et al. 1987).

Ground-water discharges along the Hanford Reach of the Columbia River have been monitored since the mid-1960s. Springs in the vicinity of the 300-Area retention basin and sewage leaching trenches were routinely sampled and analyzed for various biological, chemical, and radiological parameters. Springs along the 100-N Area, resulting from liquid waste disposal practices, have been, and are today, monitored routinely (Rokkan 1988).

In addition, special studies of ground-water discharges have been conducted periodically. The latest of these special studies was conducted in 1982 and 1983 (McCormack and Carlile 1984). This study covered approximately 41 miles of Hanford Site shoreline, identifying 115 springs or seepage areas. During this study, the constituents used as indicators of contaminated ground water were tritium and nitrate because of their predominance in much of the Hanford ground water. Uranium analyses were substituted in place of tritium on samples collected near the 300 Area where uranium is a primary constituent in the ground water. In addition to these analyses, a few samples were selected for the analysis of 90 Sr, 99 Tc, and gross beta. The McCormack and Carlile (1984) study confirmed areas of contaminated ground-water entry into the Columbia River and documented contaminant concentrations similar to the local ground water in the riverbank spring water.

Numerous attempts to sample the riverbank springs have been made since the McCormack and Carlile (1984) study. A cooperative sampling program was initiated in 1984 that concentrated its effort on the Columbia River and the ground-water seepage entering the river along the Hanford Reach. A few springs have been sampled consistently over the years as a result of this program. This program is an ongoing effort presently involving the state of

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Washington, the state of Oregon, the U.S. Environmental Protection Agency (EPA), the Washington Public Power Supply System, SEARCH, Inc., and the DOE, represented by PNL. When and where sample collection was successful, sample results obtained as a result of this program have generally been in good agreement with previous observations and provide additional background information concerning the concentrations of various radionuclides in the ground-water discharges (Jaquish et al. 1987).

SPRING CHARACTERIZATION STUDY DESCRIPTION

The FY 1988 Project Management Plan (PMP) for the SEMP included a special study to characterize the riverbank springs along the Hanford Site shoreline. The objectives of the Spring Characterization Study were to monitor the discharges along the Hanford Reach shoreline for radiological and nonradiological constituents and define points suitable for the establishment of permanent, routine spring sampling locations.

Identification of previously documented spring locations was not included in the scope of this study. Also excluded from the scope of the Spring Characterization Study, as was the case in the McCormack and Carlile (1984) study, were investigations of ground-water discharges as a function of time or as related to the flow rate of the Columbia River. No attempt was made to quantify the amount of ground water entering the river via the springs during this effort. The primary elements of the Spring Characterization Study were

- follow-up and expansion of the McCormack and Carlile (1984) study
- screening of radiologically contaminated ground-water plumes for nonradiological parameters
- · identification of permanent sampling locations
- identification of future needs relating to the ground-water/river system.

Past studies have provided significant background information concerning the concentrations of certain contaminants entering the river through the discharge of contaminated ground water. The current effort was intended to follow-up and expand on the information previously obtained. As such, sample location and analysis combinations were selected to allow for comparisons of present radionuclide concentrations with those observed in the past. Similar locations and analysis were included to determine if significant increases or decreases in contaminant concentrations were apparent. Other location/ analysis combinations were requested to determine what other, if any, contaminants were entering the river through ground water and to define the extent and location of the ground-water plumes entering the river.

Also included in this study were the sampling and analysis of irrigation returns and springs along the Franklin County shoreline to identify potentially significant sources of contaminants not associated with Hanford operations entering the river along the Hanford Reach.

A notable change in this study versus the previous spring study is the expansion of analysis to include an extensive number of nonradiological parameters. This provides much-needed background information regarding the discharge of nonradiological contaminants into the river that can be used by others in future ground- and surface-water monitoring activities as well as in the development and implementation of Resource Conservation and Recovery Act (RCRA) feasibility investigation/corrective measures studies (RFI/CMS) and Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) remedial investigations and feasibility studies (RI/FS). The inclusion of nonradiological analysis also provides some verification of observations and predictions of the ground-water monitoring projects relative to contaminants in ground water near the Columbia River shoreline.

Specific sample locations will be identified as part of this task and incorporated into the routine SEMP sampling schedule. In the future, information on specific ground-water plumes will be obtained from a specific spring or springs over time to provide time-series information relative to both the flow of the spring and the contaminant concentrations in the spring water when the discharge is active. Routine monitoring of the ground-water discharges into the Columbia River will also provide a degree of public assurance in the surveillance activities in that a visible, albeit small, transport pathway is not being ignored.

The final element of this task is intended to identify future needs or activities that would further enhance our understanding of the ground-water discharges, the interaction between these and the river, and the transport and fate of the contaminants entering the river via this pathway. In addition, better understanding of the ground-water/spring/river interactions may lead to improved river monitoring system design and allow for more complete and accurate data evaluation and interpretation of current river monitoring results.

SAMPLING PLAN

Figure 4 illustrates the extent of the study area. Hanford River Miles (HRMs) shown on this figure are approximate, starting from the Vernita Bridge and progressing 44 miles downstream at approximately 1-mile intervals to a point about 1 mile downstream of the 300 Area. HRM markers have been placed along the river, providing a field reference point for river-related activities and future relocation of specific springs.

Table A.1. Appendix A. provides the spring sample locations identified for the FY 1988 Spring Characterization Study. The locations are defined by HRM, the operational area being monitored, and the spring number, as defined in the McCormack and Carlile (1984) study. Locations were selected to be consistent, to the extent possible, with the McCormack and Carlile (1984) study to allow for meaningful comparisons and to maximize the opportunity to relocate springs with a flow adequate for the collection of a sample. The selected sample locations included springs identified as having "moderate" or "good" flows in the McCormack and Carlile (1984) study at locations just upstream, within, or just downstream of areas with elevated contaminant concentrations. Experience has shown that sampling attempts are not always successful. Therefore, multiple sites (backup sampling locations) along each operating area or point of contaminated ground-water entry were identified in an effort to obtain at least one meaningful sample from each area. Sampling activities were scheduled to coincide with low river flows, to the extent possible, to maximize the chances of finding the springs flowing and obtaining a sample.

In addition, those sites at which near-shore river water samples were scheduled are identified in Table A.l. Near-shore river water samples were collected within 5 feet of the shoreline and immediately downstream of the actively flowing springs. As such, results of near-shore river water samples provide information concerning the localized influence of the seepage and are not representative of average river concentrations. River water samples were also collected from the SEMP Columbia River monitoring stations located at Priest Rapids Dam and the Richland Pumphouse. These samples were obtained from the routine sampling systems.

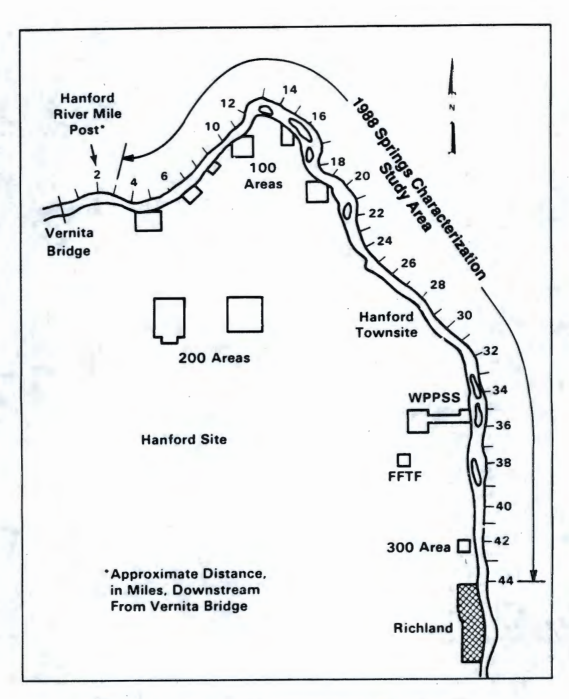


FIGURE 4. 1988 Springs Characterization Study Area

Water sample locations were also identified on the Franklin County side of the Columbia River (opposite Hanford). These samples were collected from two irrigation canal returns and from a spring entering the river as a result of extensive irrigation practices east of the river. Results of these samples provide insight on potential sources of contaminants entering the river not associated with Hanford operations, and allow for better interpretation of results obtained from river samples collected downstream.

SAMPLE COLLECTION AND ANALYSIS

Sample collection methods were consistent at all locations except for the need to "improve" those sites where the spring flow was very low and/or extremely shallow. These sites were deepened to allow for the collection of the spring water without disturbing the sediments. All suspended material was allowed to clear from the modified springs before collection of the sample. Care was taken to ensure the spring flow was not interrupted, reversed, or otherwise modified in such a manner that could influence the sample results.

A small hand suction pump was used to transfer the water from the springs to the sample containers. The pump was thoroughly rinsed before and following sample collection at each site. In addition, the pump was flushed with spring water at each site before sample collection to avoid cross contamination between samples. Flushing the pump also served to purge the spring sampling area, providing fresh spring water for sample collection. Use of the hand pump eliminated the disturbance and inadvertent collection of sediment material and minimized or eliminated aeration of the sample during the collection process, which is critical in the sampling and analysis of numerous nonradiological parameters.

Samples of river water, where specified, were collected immediately downstream of the spring entry point using methods similar to those described for the collection of spring samples. Water was collected within 2 to 5 feet of the shoreline at approximately mid-depth. Care was taken to avoid stagnant areas along the shoreline to the extent possible and collect the near-shore river samples in flowing water.

The analyses performed on each sample are identified in Table A.1, Appendix A. As with the selection of sample locations, analyses were chosen to allow for comparison with previous results on spring samples and local ground-water monitoring results. In addition, analyses were expanded at several locations to determine what additional contaminants are present in the spring water entering the river and provide background information concerning the concentrations of these contaminants. Radiological analyses are listed as a specific analysis except in the case of the gamma scan, which provides concentrations of a number of gamma-emitting radionuclides including 60Co, 106Ru, 125Sb, 134Cs, and 137Cs, as well as several others. At a minimum, all spring water samples received analysis for gross alpha, gross beta, gamma scan, tritium, and nitrate (included in the 735 and 9905 codes).

Nonradiological analyses are listed as laboratory codes (735 and 9905) in Table A.1, Appendix A. The 735 code provides for the determination of common anions such as chloride, fluoride, nitrate, phosphate, and sulphate using ion chromatography (IC). The 9905 code is made up of an extensive set of lists for various types of analyses. The 9905 code includes the dangerous waste constituents as identified by the state of Washington in WAC 173-303-9905 (Washington State Department of Ecology 1986). Analysis of the entire 9905 list provides a screening mechanism on the spring discharges, ensuring that potential contaminants are not being overlooked. The 9905 code includes analysis for the ICP metals, enhanced thiourea, enhanced pesticides, volatile organic compounds, phosphorous pesticides, direct aqueous injection analysis, enhanced herbicides, IC Report, PCB analysis, arsenic, mercury, selenium, thallium, lead, total organic carbon (TOC), cyanide, perchlorate, sulfide, ammonium ion, ethylene glycol, citrus red, total carbon (TC), and total organic halogen (TOX). Table A.2, Appendix A, lists the specific nonradiological parameters included in the 9905 (including 735) analytical code. As is evident in Table A.2, Appendix A, those samples for which the 9905 list was requested received an extensive screening for nonradiological contaminants. Because of the relatively high cost of analysis for the entire 9905 list, the number of samples analyzed for the entire list was minimal, and locations were selected in areas of known contamination to maximize the amount of useful information obtained.

All sample analyses, radiological and nonradiological, were performed by United States Testing Company (UST). All analytical procedures were the same as those used for the routine SEMP and GWPMP samples. An extensive Quality Assurance (QA) program is required for the SEMP and GWPMP that documents typical lab performance for the analysis performed (Jaquish and Bryce 1989). Detection levels for the various nonradiological constituents are included in Table A.2, Appendix A.

The type of sample container varied widely, depending on the analyses to be performed. Multiple sample containers were required at all sample locations. The bottle size, bottle type, method of sample preservation, and special handling requirements were dependent on the specific analysis performed. Table A.3, Appendix A, provides the type of sample container and sample preservation requirements for each group of analysis.

Nonradiological samples were stored in ice-filled coolers, as appropriate, and transported to the laboratory as soon as possible following sample collection. Samples to be analyzed for radiological constituents, while not having special preservation or handling requirements, were also transported to the laboratory as soon as practical following sample collection. In all cases, samples were delivered to the lab the same day they were collected.

RESULTS AND DISCUSSION

Samples were collected from 18 springs between HRM 2 and HRM 44 during 1988 as part of the Spring Characterization Study. Samples were obtained from the river along the 100-B, 100-N, 100-D, 100-H, and 300 Areas. In addition, springs were sampled that are associated with the contaminated ground-water plume originating beneath the 200 Areas, which enters the river between approximately HRM 27 (Hanford Townsite) and HRM 40. Samples were also collected upstream of all past operating areas near HRM 2.5 and downstream of all facilities at approximately HRM 43.8.

Samples were not obtained at all sites identified in the sampling plan. Locations successfully sampled during 1988 are identified in Table B.1, Appendix B. Attempts to collect samples along the 100-K and 100-F Areas were unsuccessful. In addition, several back-up sites were not sampled. A minimum of three sampling attempts were made at each specific site. River flow rates during these attempts were at levels conducive to active spring flow. This is supported by the fact that nearby springs were actively flowing and successfully sampled during the same day, sometimes minutes apart. This may be a result of changing flow patterns in the local ground water or perhaps reflect seasonal variances in flow patterns similar to those identified in the 300 Area (Schalla et al. 1988). Table B.2, Appendix B, provides the river flow rates at Priest Rapids Dam during the periods of sample collection. Daily average flow rates are provided for 2 days before sampling, the day of sample collection, and the day following sampling.

In addition to sampling the springs themselves, near-shore river water samples were collected at actively flowing spring locations along the 100-N, 100-H, and 300 Areas as well as along the Hanford Townsite. As previously discussed, these samples were intended to maximize the localized influence of the seepage and are not intended to be representative of the average river conditions. River water samples were also collected as planned from the routine SEMP Columbia River monitoring stations located at Priest Rapids Dam and the Richland Pumphouse. These samples were obtained from the routine water sampling system intakes to provide some indication of the concentrations present in the river at these locations during the spring sampling. As

scheduled, water samples were also collected from two irrigation water return canals and a seepage area on the Franklin County shoreline along the Hanford Site.

As indicated in the initial sampling plan (Table A.1, Appendix A), all the spring samples collected were analyzed for gross alpha, gross beta, gamma scan, tritium, and nitrate. Selected samples received additional radiochemical analyses depending on their locations. Individual samples from the 100-N, 100-H, and 300 Areas as well as from the Hanford Townsite area (200-Areas ground-water plume) received the extensive 9905 screening list of nonradiological analyses in addition to the radiological analysis. In addition, Columbia River water samples collected at Priest Rapids Dam and the Richland Pumphouse were analyzed for the entire list of nonradiological constituents. Similarly, the spring and irrigation return water samples collected from the Franklin County side of the river also received these analyses.

Background river concentrations are of interest in evaluating the effect of the discharge of ground water into the river along the Hanford Site. Background river concentrations are defined as those levels observed upstream of Hanford at Priest Rapids Dam or Vernita Bridge. Tables B.3 and B.4, Appendix B, provide background river concentrations for several radiological and nonradiological parameters. For those constituents where data were not available, values observed in the Priest Rapids Dam river water sample collected as part of this study are used.

Background concentrations for several contaminants in ground water have been estimated and are presented in Table B.5, Appendix B (Evans et al. 1989a and 1989b). The estimates of background contaminant concentrations were based on samples collected from areas not affected by Hanford operations. Since the spring water is ground water emanating from the riverbank and flowing into the river, ground-water background concentrations are appropriate for comparison. In some cases, background concentrations have apparently not been estimated, making comparisons with the spring sampling data difficult.

Radiological and nonradiological analytical results are presented in Tables B.6 and B.7, Appendix B, respectively. In general, the concentrations of those constituents for which previous data exist were similar during 1988 to those previously documented. In addition, contaminant concentrations found in the spring water were generally at or below those known to exist in the local ground water. Contaminant concentrations in near-shore river water samples were indicative of the localized effect of the discharge of contaminated ground water into the river. Specific results are discussed in more detail in the following subsections.

100-B AREA

The 100-B Area has not been identified as an area of extensive ground-water discharge to the Columbia River, although several springs have been documented along the shoreline that contained elevated levels of tritium and nitrate. Only one active spring was found in the vicinity of the 100-B Area during the 1988 sampling activities. This spring was located just upstream of the 181-B Water Intake at about HRM 3.75. Spring locations downstream of the water intake, identified in previous studies, were visited on several occasions; however, no flow was present. Tritium and nitrate concentrations observed during 1988 in the spring water (1100 pCi/L and 6700 ppb, respectively) were similar to those observed in 1982 and slightly lower than those present in the local ground water during 1988 (McCormack and Carlile 1984; Evans et al. 1989a and 1989b).

100-K AREA

As in the case of the 100-B Area, the 100-K Area is not considered a major source of ground-water discharge to the Columbia River. Several spring locations were identified during 1982 shoreline inspections (McCormack and Carlile 1984). No spring flow was observed during any of the three attempts to collect samples from the 100-K Area.

100-N AREA

The 100-N Area shoreline has been identified as a major ground-water discharge area containing elevated levels of several radionuclides and nitrate. The N8T Monitoring Well, operated by Westinghouse Hanford Company, provides a reliable method for the collection of ground water that is representative of the spring water under most conditions. This well is located very near the river and is sampled routinely by Westinghouse Hanford Company as a record of the discharge via the springs from the 100-N Area (Rokkan 1988). Active springs were not observed in the immediate vicinity of the monitoring well during the sampling efforts, although they are known to flow periodically in this region. Large boulders placed along the shoreline hinder the location and sampling of springs that may be present in the area. An additional spring was located and sampled downstream of the riprap. This spring has shown increased flows since the closure of the 1301-N Liquid Waste Disposal Facility (LWDF) and startup of the 1325-N LWDF.

Radionuclide concentrations in the 100-N Area springs were similar to those observed in the past and were indicative of the local ground water. Table 1 presents the radionuclide concentrations observed in the springs and the 100-N Area ground water during 1988. Gross beta, tritium, 60co. 90sr. and 125Sb are present at elevated levels in the spring water, well above background river water concentrations (Table B.3); however, the concentrations are within the range of values observed in the local ground water. Of these, gross beta, tritium, and 90Sr are above the applicable drinking water standard (DWS). The 90Sr concentration is above the DOE Derived Concentration Guide. Gross beta and 90Sr discharges to the river via the 1301-N LWDF and associated springs have decreased during recent years, primarily as a result of the smaller volumes of water being discharged into the LWDF (Rokkan 1988). However, as shown in Table 2, the concentrations of these constituents in the N8T Monitoring Well water have not shown a decrease. This is probably due to the inventory of 90Sr remaining in the soil column as a result of past effluent disposal practices.

Extensive nonradiological analyses were performed on the sample collected from the N8T Monitoring Well. In most instances, nonradiological

TABLE 1. Contaminant Concentrations in Spring and Ground Water Along the 100-N Area Shoreline During 1988

	-		•	
		Concentr	ation, pCi/L	
	(-)		Spring Water	
<u>Radionuclide</u>	Ground Water(a)	Date	HRM 8.9	HRM 9.5
Gross Alpha	3.07 ± 2.38	9/06/88	0.55 ± 0.61	0.37 ± 0.46
		9/13/88	0.0004 ± 0.28	NS(b)
Gross Beta	$40,800 \pm 1,110$	9/06/88	$13,800 \pm 100$	74 ± 8
		9/15/88	$10,800 \pm 296$	NS
Tritium	$459,000 \pm 2,210$	9/06/88	$74,000 \pm 700$	$111,000 \pm 870$
		9/15/88	$75,800 \pm 908$	NS
90 _{Sr}	$13,800 \pm 262$	9/06/88	$6,680 \pm 260$	0.31 ± 0.09
		9/15/88	$7,270 \pm 192$	NS
60 _{Co}	128 ± 24	9/06/88	45.0 ± 4.5	30.0 ± 4.0
		9/15/88	53.4 ± 19.4	NS
106 _{Ru}	116 ± 67.3	9/06/88	8.9 ± 11.4	13.5 ± 14.6
		9/15/88	12.6 ± 71.2	NS
125 _{Sb}	108.00 ± 47.50	9/06/88	43.1 ± 6.4	, ND(c)
		9/15/88	ND ND	NS
137 _{Cs}	6.01 ± 4.79	9/06/88	-0.2 ± 1.0	0.4 ± 1.1
		9/15/88	-0.4 ± 4.4	NS
NO ₃	67,100 ppb	9/06/88	NA(d)	NA
		9/15/88	28,630 ppb	NS

⁽a) Maximum reported value from any one well in the general area during 1988.

TABLE 2. Selected Contaminants in 100-N Area Spring Water 1986 through 1988

	a)				
Year	Beta	3 _H	60 _{Co}	90 _{Sr}	NO3
1986	$9,120 \pm 610$	$65,100 \pm 800$	68 ± 10	$6,060 \pm 370$	9.5 mg/L
1987	$10,500 \pm 350$	$84,300 \pm 1,120$	98 ± 30	$5,830 \pm 380$	-
1988	$13,800 \pm 110$	$74,000 \pm 730$	45 ± 5	$6,680 \pm 260$	-

⁽a) The maximum concentration is reported for those years having more than one sample collected.

⁽b) NS = Not sampled.

⁽c) ND = Not detected.

⁽d) NA = Not analyzed.

contaminant concentrations were below detection levels. When detectable, contaminant concentrations were generally at or below the estimated background levels for Kanford ground water. Nitrates were observed at elevated levels, consistent with past ground-water monitoring and spring sampling results. The concentration of nitrate in the spring samples was well above those typical of the river water, but below the DWS in all cases.

100-D AREA

The 100-D Area has not been identified as a major ground-water discharge source directly to the river. Predominant ground-water flow in this region is to the east-northeast, across the northernmost tip of the Hanford Site. One spring was identified along the 100-D Area at HRM 11 during the McCormack and Carlile (1984) study. This spring was located and successfully sampled during 1988. Concentrations of tritium and NO3 found in 1988 were similar to those observed in 1982 (McCormack and Carlile 1984). The concentration of tritium was below the analytical detection level, well below the DWS of 20,000 pCi/L. Nitrate levels were also well below the DWS and lower than those observed in the nearby ground water. The concentration of 90Sr was elevated with respect to river levels. However, it remained below the 8 pCi/L DWS and was lower than the maximum observation made in the local ground water during the first half of 1988 (Evans et al. 1989b).

100-H AREA

Springs identified along the 100-H Area during the 1982 shoreline seepage investigation were generally not flowing during the 1988 sampling attempts. Samples were successfully obtained on two occasions from a small spring at HRM 15. The first sample was collected during July and the second in September. Results from the two samples varied considerably, particularly in the case of tritium. This is believed to be a result of the influence of river water on the spring water composition during the second sampling. It was apparent that the river level had been rather high a short time before sampling. Therefore, it is probable that the water sampled was a mixture of river water and ground water.

Results of spring and ground-water samples collected along the 100-H Area during 1988 are included in Table 3. Tritium concentrations in the springs during the first sampling ($2520 \pm 220 \text{ pCi/L}$) were considerably higher than those found later in the year ($550 \pm 180 \text{ pCi/L}$). Nitrate levels were also lower in the September sample than they were in July (less than 500 ppb and 2700 ppb, respectively), similar to those observed in the river, further supporting the idea that the sample in September was composed primarily of river water. These concentrations were, however, similar to those observed in nearby springs during earlier studies and below those typically observed in the local ground water (McCormack and Carlile 1984; Evans et al. 1989a and 1989b).

Results of the extensive nonradiological analysis performed on the sample collected during September revealed concentrations similar to back ground levels in those few cases where the constituents were above the analytical detection level. The 100-H Area is known to have elevated levels

TABLE 3. Selected Radionuclide Concentrations in Spring and Ground Water Along the 100-H Shoreline During 1988

	C	oncentration, pCi/L	
	(a)	Spring Water	HRM 15.0
Radionuclide	Ground Water(a)	9-12-88	7-11-88 NA(b)
Gross Alpha	1230	0.30 ± 0.35	NA
Gross Beta	1920	1.63 ± 1.27	NA
3H	5550	545 <u>+</u> 178	2520 ± 220
60 _{Co}	6.08	-0.55 ± 1.92	0.9 ± 1.8
90sr	10.30	NA	0.2 ± 0.1
99 _{Tc}	3835	NA	NA
106 _{Ru}	NR(c)	-1.69 ± 20.1	17 ± 15
125 _{Sb}	NR	NR	NR
137 _{Cs}	NR	-0.94 ± 1.62	-1.1 ± 1.7
U Total	166	0.12 ± 0.06	NA

⁽a) Maximum reported value from any one well in the general area during 1988. Uncertainty terms associated with results not reported.

⁽b) NA = Not analyzed.

⁽c) NR = Not reported.

(above DWS) of chromium in the ground water (Liikalla 1988); however, no evidence of chromium was detected in the spring sample. The apparent presence of river water in the spring water at the time of sampling may have diluted the chromium to less than the detectable level. In addition, examination of the chromium plume in the 100-H Area ground water indicates that the chromium may be entering the river downstream of the spring that was sampled during 1988. Inspection of the shoreline downstream of HRM 15 in areas of previously identified seepage failed to locate any active springs during the 1988 sampling attempts.

100-F AREA

No active seepage areas were located during several shoreline inspections along the 100-F Area. Two springs were documented along the 100-F Area (HRM 18.5 to HRM 20) during earlier investigations (McCormack and Carlile 1984). Three attempts to locate these springs in these areas during 1988 were unsuccessful.

300 AREA

Seepage of the contaminated 300-Area ground water is of particular interest because of its proximity to drinking water supply intakes. The 300-Area water system intake is located directly downstream of active seepage areas and within the contaminated ground-water plume as identified in recent ground-water monitoring reports (Schalla et al. 1988; Jaquish and Bryce 1989). In addition, the city of Richland withdraws river water for its water supply approximately 5 miles downstream of the 300 Area. Richland is the nearest community downstream of Hanford using the Columbia River as its drinking water source.

The 300 Area is also the focus of ongoing Hazardous Waste Ground-Water Monitoring Compliance Program characterization studies. Initiated in 1985 and expanded in 1986, this effort has provided an extensive amount of information relative to the hydrogeology and contaminant history of the 300 Area. Most recently, PNL-6716, Interim Characterization Report for the 300 Area Process Trenches (Schalla et al. 1988), was issued, providing the basis

for comparisons of the concentrations observed in the riverbank spring water with the local ground water.

Table 4 presents the radionuclide concentrations in spring and ground water along the 300 Area shoreline during 1988. The concentrations were similar to those found during earlier years. As expected, the gross alpha, gross beta, 234 U, 235 U, and 238 U concentrations were elevated with respect to river concentrations and similar to those reported in the ground water beneath the 300 Area (Schalla et al. 1988). Concentrations of other radioisotopes were below the analytical detection levels.

One of the springs along the 300 Area (Spring 42-2, HRM 42.1) has been sampled routinely over the years as part of the cooperative radiochemical laboratory intercomparison effort previously discussed. Results of samples collected from Spring 42-2 during recent years have shown relatively good agreement and are comparable with those observed in the local ground water.

TABLE 4. Selected Radionuclide Concentrations in Spring and Ground Water Along the 300 Area Shoreline During 1988

		C	oncentrat	ion, pCi/L	
Radionuclide	Ground Wa	iter(a)	HRM	Spring 42.1	HRM 42.3
Gross Alpha		7.6	8.0	± 1.5	6.5 ± 1.4
Gross Beta	67.6	7.69	11.6	± 3.3	9.8 ± 2.2
Tritium	5940	291	346	± 172	283 ± 170
60 _{Co}	7.68 ±	7.05	0.25	± 0.80	1.6 ± 2.2
90 _{Sr}	7.29 +	1.54	0.16	± 0.07	NA(b)
99 _{Tc}	214	2.9		NA	NA
137 _{Cs}	8.25	6.37	0.40	± 0.60	-0.36 ± 2.08
234 _U	21.2	0.4	4.5	± 0.2	3.48 ± 0.18
235 _U	0.84	0.08	0.36	± 0.06	0.24 ± 0.05
238 _U	20.8	0.4	4.6	± 0.2	3.00 ± 0.17
U Total	101	5	9.5	<u>+</u> 0.4	6.7 ± 0.4

⁽a) Maximum reported value from any one well in the general area during 1988.

⁽b) NA = Not analyzed.

Spring 42-2 samples were analyzed for nonradiological pollutants during the 1988 study. Results of this sampling revealed elevated concentrations (with respect to background ground-water concentrations) of chloroform, copper, nitrate, and zinc in the spring water. All of these contaminants have been identified as being present in the local ground water (Schalla et al. 1988). Table 5 provides the concentrations of these constituents in the spring water and in the local ground water during 1988. Nitrates were at concentrations similar to those previously experienced in the springs and known to exist in the ground water. The concentrations of copper and zinc were well within the range reported in the local ground water. Fluoride, present in the ground water at elevated levels in some locations, was not detected in the spring water samples.

Chloroform has been documented to be present in the ground water beneath the 300 Area (Schalla et al. 1988). The concentrations of chloroform observed in the spring water during 1988 were similar to those found in the local ground water (Schalla et al. 1988; Evans et al. 1989a and 1989b). Concentrations of other organic contaminants identified in the ground water, such as methylene chloride, PCE, TCE, 1,1,1-trichloroethane, and trans-1,2-dichloroethylene were not detected in the 1988 spring samples. Of these, methylene chloride, PCE, TCE, and 1,1,1-trichloroethane have been only sporadically detected in a few locations and would not be expected at detectable levels in the spring water. Trans-1,2-dichloroethylene,

TABLE 5. Selected Nonradiological Contaminants in Spring and Ground Water Along the 300 Area Shoreline

	Concentration, ppb				
		Spring Water			
Contaminant	Ground Water	HRM 42.1	HRM 42.3		
NO3	66,800	1,699	9,183		
Chloroform	42	24	19		
Zinc	232	23	10		
Copper	70	34	<10		

⁽a) Maximum reported value from any one well in the general area during 1988.

consistently found at relatively high levels, has been detected primarily in the intermediate and deep wells (Schalla et al. 1988). Consequently, this contaminant may be entering the river through submerged seepage, consistent with typical ground-water flow patterns (Raymond and Brown 1963).

HANFORD TOWNSITE (200-AREAS GROUND-WATER PLUME)

The 200-Areas contaminated ground-water plume has long been identified as generally flowing from the 200 Areas in a easterly/southeasterly direction, discharging into the Columbia River in the vicinity and downstream of the old Hanford Townsite (Figures 2 and 3). Primary contaminants present in the ground water near the river are tritium, 129 I, and nitrate (Jaquish and Bryce 1989). Technetium-99 was also identified in springs sampled along this stretch of the river (McCormack and Carlile 1984). Other studies have implicated the possible presence of 60 Co and 90 Sr in the ground water along the Hanford Townsite (Buske and Josephson 1989). Analytical methods used in the routine ground-water monitoring program and for special SEMP samples typically are not sensitive enough to detect these radionuclides at their extremely low concentrations.

Table 6 contains the concentrations of selected radionuclides in the spring and ground water along the old Hanford Townsite during 1988. Radionuclide concentrations observed during the year were generally similar to those previously reported (McCormack and Carlile 1984). Radionuclide concentrations in the springs were also comparable with those in the nearby ground water (Evans et al. 1989a and 1989b). It appears that the highest concentrations in the plume discharge zone remain at approximately HRM 28.

A notable difference between earlier investigations and the present study was the southern extent of the contaminated springs associated with the contaminated 200-Areas ground-water plume. Consistent with the findings of the GWPMP, the southern extent of the plume reaches nearly to the northern border of the 300 Area. Table 7 lists the tritium concentrations in spring water from the northern edge of the 200-Areas ground-water plume to the southern boundary of the Hanford Site (also see Figure 5).

Selected Radionuclide Concentrations in Spring and Ground Water Along the Hanford Townsite During 1988 TABLE 6.

		Concentration, pCi/L						
	4-5		Sprin	Water				
Radionuclide	Ground Water (a)	HRM 25.75	HRM 27.25	HRM 27.5	HRM 28.1	HRM 28.5		
Gross Alpha	4.1 ± 0.9	NA (b)	2.5 ± 2.1	2.1 ± 1.0	. 2.3 ± 1.1	1.5 ± 0.9		
Gross Beta	69.9 ± 5.78	NA	4.3 ± 4.1	14.2 ± 2.7	48 ± 5	45 ± 5		
3,	246,000 ± 1,610	264 ± 169	7,420 ± 296	72,000 ± 890	155,000 ± 1,290	145,000 ± 1,250		
60 _{Co}	14.7 ± 8.15	-0.08 ± 1.2	NA	1.1 ± 3.7	4.7 ± 4.8	2.8 ± 3.2		
90 _{Sr}	MR (C)	NA	NA	-0.07 ± 0.33	0.07 ± 0.3	0.0014 ± 0.3		
99 _{TC}	350 ± 4	NA	NA	48 ± 2	223 ± 3	215 ± 3		
137 _{Cs}	NR	0.4 ± 1.4	NA	0.6 ± 2.8	-3.7 ± 3.3	-1.3 ± 2.6		

Maximum reported value from any one well in the general area during 1988. NA = Not analyzed. NR = Not reported.

Tritium Concentration in Spring Water Along the Hanford Shoreline from HRM 25.75 to HRM 44 During 1988 TABLE 7.

HRM	Tritium Concentration. pCi/L
25.75	260 ± 170
27.25	$7,420 \pm 300$
27.5	72,000 <u>+</u> 890
28.1	$155,000 \pm 1,290$
28.5	$145,000 \pm 1,250$
38.25	2,630 ± 230
38.8	682 <u>+</u> 180
41.75	6,580 ± 310
42.0	1,070 <u>+</u> 190
42.1	346 ± 170
42.3	283 ± 170
43.6	65 ± 160
	25.75 27.25 27.5 28.1 28.5 38.25 38.8 41.75 42.0 42.1 42.3

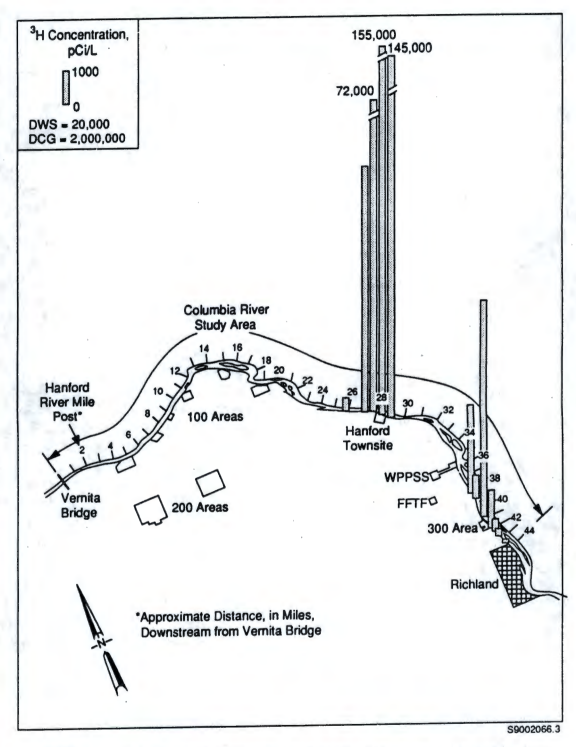


FIGURE 5. Tritium Concentrations in Spring Water Along the Hanford Shoreline from HRM 25.75 to HRM 44 During 1988

COLUMBIA RIVER

Samples of the Columbia River were collected as part of the Spring Characterization Study for comparison and to provide information relative to the impact of the ground-water discharge. Grab samples were collected from the routine sample locations using the routine sampling system intakes. In addition, river water samples were collected immediately downstream of selected riverbank springs in order to identify the localized influence of the seepage on the river. The concentrations observed in these near-shore river samples are not representative of average river water concentrations and are indicative of the elevated concentrations attributable to the discharge of contaminated ground water into the river.

Radiological and nonradiological sample results from Priest Rapids Dam and the Richland Pumphouse, observed above the detection level, are included in Tables 8 and 9, respectively. As was the case for selected springs, samples from these two locations received the extensive 9905 list of nonradiological analysis. As expected, most contaminants were not detectable in the river water samples. Radiologically, only the concentrations of tritium, 99Tc, 234U, and 238U were above the analytical detection levels. The tritium and 99Tc concentrations were higher at the Richland Pumphouse than at Priest Rapids Dam, although the uncertainties associated with the 99Tc results overlapped on the two locations. The tritium concentration was considerably higher at the Richland Pumphouse location, well above the 1988 average concentration of 130 pCi/L (Jaquish and Bryce 1989). This is expected since the annual average is determined using composite samples that flatten out the extremes during the year and because the grab sample in this study was obtained during a low flow condition that would maximize the influence of the ground-water discharges to the river. Nonradiological sample results (Table 9) were similar at the two locations and similar to concentrations previously reported (McGavock 1987).

Table 10 lists the tritium and nitrate results of the near-shore river water samples collected immediately downstream of specific riverbank springs. Also included in this table are the corresponding concentrations observed in

TABLE 8. Radiological Results of Water Samples Collected at Priest Rapids Dam and the Richland Pumphouse on September 13, 1988

	Concentration, pCi/L				
Constituent	Priest Rapids Dam	Richland Pumphouse			
Tritium	180 ± 170	580 <u>+</u> 180			
99 _{Tc}	0.6 ± 1.2	1.8 ± 1.2			
234 _U	0.16 ± 0.04	0.18 ± 0.04			
238 _U	0.13 ± 0.04	0.16 ± 0.04			

TABLE 9. Nonradiological Sample Results of Water Samples Collected at Priest Rapids Dam and the Richland Pumphouse on September 13, 1988

	Concentration, ppb					
Constituent	Priest Rapids Dam	Richland Pumphouse				
Strontium	125	120				
Zinc	12	9				
Calcium	21,657	21,362				
Barium	33	32				
Sodium	2,452	2,563				
Manganese	14	8				
Potassium	811	862				
Iron	160	37				
Magnesium	4,777	4,567				
Nitrate	<500	554				
Sulphate	10,336	10,802				
Chloride	895	1,019				

the spring water itself. Concentrations of radiological constituents in the near-shore river water samples were generally similar to those observed in average river water samples, typically below the analytical detection level. Exceptions to this included tritium in samples collected at various locations along the Hanford Reach, 90 Sr concentrations in the near-shore river sample near the N-spring, and uranium concentrations in the near-shore river samples collected at the 300 Area, which were found at levels similar to the local ground water.

TABLE 10. Tritium and Nitrate in Near-Shore River and Spring Samples Along the Hanford Site During 1988

Concentration			
Sample Type	Tritium, pCi/L	Nitrate, ppb	
Spring	$75,800 \pm 910$	28,630	
River	$76,400 \pm 910$	NA(a)	
Spring	545 ± 180	<500	
River	70 ± 130	<500	
Spring	$72,000 \pm 890$	12,713	
River	$26,400 \pm 525$	NA	
Spring	$155,000 \pm 1,290$	31,040	
River	$158,000 \pm 1,250$	31,290	
Spring	346 ± 170	2,149	
River	485 ± 180	1,697	
	Spring River Spring River Spring River Spring River Spring River Spring	Sample Tritium, pCi/L Type Tritium, pCi/L Spring 75,800 ± 910 River 76,400 ± 910 Spring 545 ± 180 River 70 ± 130 Spring 72,000 ± 890 River 26,400 ± 525 Spring 155,000 ± 1,290 River 158,000 ± 1,250 Spring 346 ± 170	

⁽a) NA = Not analyzed.

Contaminant concentrations in the near-shore river samples reflect the localized effects of the ground-water discharge to the river. Specific sample locations were chosen to maximize the potential to detect the influence of the springs on the river concentrations. As expected, the radionuclide concentrations found in the near-shore river samples were generally elevated with respect to the average concentrations in the river as defined by the samples collected at Priest Rapids Dam and the Richland Pumphouse. Most near-shore river water samples displayed concentrations between those observed in the spring and the average river water concentrations. In some instances, the concentrations observed in the near-shore samples were at or above the corresponding spring water concentrations, indicating a significant contribution from the ground water on local river water concentrations. This phenomena has been reported in previous studies (McCormack and Carlile 1984).

FRANKLIN COUNTY

An understanding of all sources of pollutants entering the river along the Hanford Site, whether or not they are associated with Hanford operations, is needed to fully understand the results of Columbia River monitoring downstream of Hanford and evaluate the subsequent impacts attributable to Hanford. Samples were collected from an area of extensive ground-water seepage across from and just above the 300 Area (HRM 41) and from two irrigation return canals that enter the river upstream of the Richland Pumphouse Columbia River monitoring station. The first canal enters the river at approximately HRM 32 at Ringold, and the second canal enters near the southern Site boundary at Byers Landing, HRM 44.

Samples collected from the Franklin County shoreline contained detectable concentrations of several constituents at levels above those known to exist in Columbia River water. Tables 11 and 12 provide the concentrations of those constituents. Interestingly, gross alpha, gross beta, 234U, 235U, and ²³⁸U concentrations were found at levels considerably higher than those

TABLE 11. Selected Radiological Contaminant Concentrations in Franklin County Seepage and Irrigation Return Canal Water During 1988

	Concentration, pCi/L					
Constituent	Ringold Irrigation	Sagemoor Seep	Byers Landing Irrigation	River Background(a)		
Gross Alpha	17 ± 0.7	2.0 ± 1.0	0.7 ± 0.5	0.3 ± 0.2		
Gross Beta	9.9 ± 2.5	2.6 ± 1.7	6.4 ± 2.1	1.0 ± 0.4		
Tritium	NA(b)	380 ± 170	NA	70 ± 6		
60 _{Co}	1.6 ± 3.5	3.0 ± 3.0	1.3 ± 2.5	<0.0002		
90 _{Sr}	0.39 ± 0.36	0.09 ± 0.35	0.42 ± 0.36	0.10 ± 0.02		
137 _{Cs}	0.9 ± 3.0	2.5 ± 3.0	0.8 ± 2.0	<0.003		
234 _U	3.0 ± 0.2	4.6 ± 0.2	2.6 ± 0.2	0.20 ± 0.03		
235 _U	0.17 ± 0.04	0.14 ± 0.03	0.15 ± 0.05	0.006 ± 0.003		
238 _U	2.3 ± 0.1	3.9 ± 0.2	2.3 ± 0.2	0.17 ± 0.02		

¹⁹⁸⁸ average at Priest Rapids Dam. (a) (b)

NA = Not analyzed.

TABLE 12. Selected Nonradiological Contaminant Concentrations in Franklin County Seepage and Irrigation Return Canal Water During 1988

	Concentration, ppb				
Constituent	Ringold Irrigation	Sagemoor Seep	Byers Landing Irrigation	River Background (a)	
Calcium	52,916	93,019	45,680	21,657	
Barium	53	64	63	33	
Sodium	47,725	72,148	42,975	2,452	
Vanadium	15	8	13	<5	
Aluminum	<150	692	<150	<150	
Manganese	11	93	10	14	
Potassium	7,316	2,605	5,067	811	
Iron	200	696	155	160	
Magnesium	25,446	48,426	21,418	4,777	
Arsenic	6	<5	5	<5	
Selenium	5	10	<5	<5	
TOC (Total Organic Carbon)	2,500	2,000	2,600	1,281	
TC (Total Carbon)	47,290	39,928	40,770	13,320	
TOX (Total Organic Halogen)	14	12	28	8.0	
Nitrate	10,795	52,320	15,631	<500	
Sulphate	63,760	238,000	50,670	10,336	
Chloride	15,810	42,370	13,560	895	

⁽a) Sample from Priest Rapids Dam Location.

observed in the river. In addition, 90Sr concentrations in the two irrigation return canals were slightly higher than those typically found in the river, although the uncertainties associated with these results were very large with respect to the results themselves.

The elevated gross alpha and beta concentrations are attributed to the corresponding elevated levels of uranium present in these water samples. Uranium is known to be present in the ground water in Franklin County (WDSHS 1988); therefore, it is not surprising for it to be present in the springs entering the river along this stretch of the river. Uranium is also known to be present in the Spokane River drainage system, which feeds into the

Columbia River upstream of Grand Coulee Dam. The irrigation water in the two canals sampled originated from behind Grand Coulee Dam, near the mouth of the Spokane River, perhaps contributing to the elevated concentrations. Another possible contributor to the uranium concentrations in the irrigation return water is phosphate fertilizer, used extensively in agricultural applications in this region. The phosphate rock ores used in fertilizer production typically contain above-average concentrations of uranium (NCRP 1984).

CONCLUSIONS

The 1988 Spring Characterization Study has confirmed that the type and concentrations of contaminants in the riverbank springs along the Hanford shoreline are within the range known to exist in the ground water near the river. In addition, the location and extent of ground-water discharge zones compare favorably with those plotted or predicted by the GWPMP in recent reports. The major contaminated ground-water discharge areas continue to be the 100-N Area, the old Hanford Townsite (200-Areas ground-water plume), and the 300 Area. The 200-Areas ground-water plume has expanded as expected and is now discharging into the river farther south than previously observed, nearly to the northern edge of the 300 Area. The ground-water flow patterns beneath the 100-N LWDFs also appear to be changing as a result of different effluent disposal practices.

Discharges of a few contaminants above the DWS into the Columbia River were observed in some areas. Discharge volumes remained very small relative to the flow of the Columbia River. However, as in the past, localized areas of impact were observed within the river near the discharge zone with radio-nuclide concentrations above the DWS. Past experience has shown these zones of influence to be rapidly dissipated within the mainstream of the river (Haney 1957). Downstream river sampling also demonstrated the impact of ground-water discharges to be very small, negligible in most cases. In all cases, river water concentrations observed outside of the immediate discharge zones were well below DWS.

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Washington State Department of Ecology (WDOE). 1986. "Dangerous Waste Regulations." In Washington Administrative Code, WAC 173-303, Washington State Department of Ecology, Olympia, Washington.

APPENDIX A

SAMPLING LOCATIONS ANALYSES INFORMATION

TABLE A.1. Scheduled Spring Sample Locations and Analyses - 1988

Area	HRM(=)	Spring No.	River	Analysis
Above 186-B	3.3	3-2		Alpha, Beta, 3H, -35(b)
1 66 -B	4.2	4-1	X	Alpha, Beta, 34, 9965(b)
100-8	5.6	5-1		Alpha, Beta, 3H, 735
Above 186-K	6.6	6-1		Alpha, Beta, 5H, 735
166-K	6.8	7-6		Ahpha, Beta, 3H, 735
Above 166-N	7.5-8.2	7-4/8-1		Alpha, Beta, 3H, 735
166-N	8.6-8.6	6-16/6-11		Alpha, Beta, SH, 96Sr, Games, 735
166-N (NST)	8.9	NST	X	Alpha, Beta, 3H, 98Sr, Gamma, 9985
166-H	9.25	9-2/9-4		Alpha, Beta, 3H, 96Sr, Games, 735
Below 166-N	9.5-10.6	MA(c)		Alpha, Beta, 3H, 96Sr, Games, 735
196-D	11.6	11-1		Alpha, Beta, 3H, 96Sr, 735
166-H	14.5	14-4/14-5		Alpha, Beta, 3H, U-iso, 735
166-H	15.6	15-6	x	Alpha, Beta, 3H, U-iso, 9965
100-H	15.5	15-5		Alpha, Bets, 3H, 735
160-F	18.6	18-6		Alpha, Beta, 3H, 735
100-F	18.6	18-2		Alpha, Bets, 3H, 9965
Hanford Ferry Landing	25.75	25-4		Alpha, Beta, 3H, 735
Hanford Townsite	27.6	27-1		Alpha, Beta, 3H, 735
Hanford Townsite	27.25	27-2		Alphs, Beta, 3H, 735
Hanford Townsite	27.5	27-3	X	Alpha, Beta, 3H, 96Sr, 99Tc, 735
Hanford Townsite	28.1	28-2	X	Alpha, Beta, 3H, 96Sr, 99Tc, 9965
Hanford Townsite	28.5	28-4		Alpha, Beta, 3H, 96Sr, 99Tc, 735
Ringold Island	31.6	31-5		Alpha, Beta, SH, 735
WPPSS Intake	35.6	35-1		Alpha, Beta, 3H, 735
Above Wooded Island	37.5	37-2		Alpha, Bets, SH, 735
Wooded Island	38.25	38-1		Alpha, Beta, 3H, 735
Wooded Island	38.8	38-16/39-6		Alpha, Beta, 3H, 735
Above 366	41.5-41.8	41-1/41-2		Alpha, beta, 3H, 735
366	42.0	42-1		Alpha, Beta, 3H, U-iso, 735
366	42.1	42-2	X	Alpha, Beta, 3H, U-iso, 238U, 9985
360	42.3	42-4		Alpha, Beta, 3H, U-iso, 236U, 9965

TABLE A.1. (contd)

Area	HRM	Sample No.	River	Analysis
Port of Benton	43.6	48-1		Alpha, Beta, 3H, 735
Byers Lndg: Irr. Return	(q)	NA		Alphs, Bets, 98Sr, U-iso, 9985
Slide Seep		NA		Alpha, Beta, SH, 985r, U-iso, 9985
Ringold Irr. Return		NA		Alpha, Beta, 96Sr, U-iso, 9965
Priest Rapids Das		NA	X	Alpha, Beta, SH, 96Sr, 99Tc, U-iso, 9985
Richland Pumphouse		NA	x	Alpha, Bets, SH, 985r, 99Tc, U-ise, 9985

⁽a) HRM = Hanford River Mile. (b) Analytical Laboratory Codes. (c) NA = Not applicable.

TABLE A.2. Nonradiological Analysis

Analytical Code	Compound	MDL (ppb)
726	ICP Metals	
A01	Beryllium	5
A03	Strontium	20
A04	Zinc	5
A05	Calcium	50
A06	Barium	6
A07	Cadmium	2
A08	Chromium	10
A10	Silver	. 10
A11	Sodium	200
A12	Nickel	10
A13	Copper	10
A14	Vanadium	5
A15	Antimony	100
A16	Aluminum	150
A17	Manganese	5
A18	Potassium	100
		30
A19 A50	Iron Magnesium	50
727	Enhanced Thiourea	
A24	Thiourea	200
A25	1-Acetyl-2-Thiourea	200
A26	1-(0-Chlorophenyl)-Thiourea	200
A27	Diethylstilbestrol	200
A28	Ethylenethiourea	200
A29	1-Naphthyl-2-Thiourea	200
A32	N-Phenylthiourea	500
729	Enhanced Pesticides	
A33	Endrin	0.1
A34	Methoxychlor	3
A35	Toxaphene	1
A36	Alpha BHC	0.1
A37	Beta BHC	0.1
A38	Gamma BHC	0.1
A39	Delta BHC	0.1
A40	DDD	0.1
A41	DDE	0.1
A42	DDT	0.1
A43	Heptachlor	0.1
A44	Heptachlor Epoxide	0.1
A44	Dieldrin	0.1
A40 A47	Aldrin	0.1
		1
A48	Chlordane	1

Analytical Code	Compound	MDL (ppb)	1
729	Enhanced Pesticides (contd)		
A52	Endosulfan II	0.1	
C62	Chlorobenzilate	30	
X10	DBC	1.0	(Expected)
731	Volatile Organic Compounds		
A61	Tetrachloromethane	5	
A62	Benzene	5	
A63	Dioxane	500	
A64	Methylethyl Ketone	10	
A65	Pyridine	500	
A66	Toluene	5	
A67	1,1,1-Trichloroethane	5 5 5 5	
A68	1,1,2-Trichloroethane	5	
A69	1,1,2-Trichloroethene	5	
A70	Perchloroethylene	5	
A71	Xylene (O, P)	5	
B14	Xylene (M)	5	
A76	Methyl Bromide	10	
A77	Carbon Disulfide	10	
A78	Chlorobenzene	10	
A79	2-Chloroethyl vinyl ether	10	
A80	Chloroform	5	
A81	Chloromethane	10	
A89	1,1-Dichloroethane	10	
A90	1,2-Dichloroethane	10	
A91	Trans-1,2-Dichloroethene	10	
A92	1,1-Dichloroethene	10	
A93	Methylene Chloride	10	
A94	1,2-Dichloropropane	10	
A95	1,3-Dichloropropenes	10	
B06	1,1,2,2-Tetrachloroethane	10	
B08	Bromoform	10	
B13	Vinyl Chloride	10	
H68	Hexone	10	
	The following have been forward searched:		
A72	Acrolein	10	
A73	Acrylonitrile	10	
A74	Bis (Chloromethyl) Ether	10	
A75	Bromo Acetone	10	
A82	Chloromethylmethylether	10	
A83	Crotonaldehyde	10	
A84	1,2-Dibromo-3-Chloropropane	10	

TABLE A.2. (contd)

Analytical Code	Compound	MDL (ppb)
731	Volatile Organic Compounds (con-	
A85	1,2-Dibromoethane	10
A86	Dibromomethane	10
A87	1,4-Dichloro-2-Butene	10
A88	Dichlorodifluoromethane	10
A96	N,N-Diethylhydrazine	10
A99	Hydrogen Sulfide	10
B01	Iodo Methane	10
B02	Methacrylonitrile	10
B03	Methanethiol	10
B04	Pentachloroethane	10
805	1,1,1,2-Tetrachloroethane	10
B09	Trichloromethanethiol	10
B10	Trichlorofluoromethane	10
B11	Trichloropropane	10
B12	1,2,3-Trichloropropane	10
B15	Diethylarsine	10
C71	Formaldehyde	500
C04	Methyl Methacrylate	10
H06	Ethyl Methacrylate	10
B19	Acetonitrile	3 ppm
H05	Ethylene Oxide	3 ppm
733	Semivolatile Organic Analysis	
B26	Aniline	10
B30	Benz(A)Anthracene	10
B33	Benzidine	10
B34	Benzo(B)Fluoranthene	10
B38	Bis(2-Chloroethoxy)Methane	10
B39	Bis(2-Chloroethyl)Ether	10
B40	Bis(2-Ethylhexyl)Phthalate	10
B41	4-Bromophenyl Phenyl Ether	10
B42	Butyl Benzyl Phthalate	10
B45	P-Chloroaniline	10
B46	P-Chloro M-Cresol	10
B48	2-Chloronaphthalene	10
B49	2-Chlorophenol	10
B50	Chrysene	10
B55	Dibenz(A, H) Anthracene	10
B60	Di-N-Butyl Phthalate	10
B61	1,2-Dichlorobenzene	10
B62	1,3-Dichlorobenzene	10
B63	1,4-Dichlorobenzene	10
B64	3,3'-Dichlorobenzidine	20
B65	2,4-Dichlorophenol	10
B67	Diethyl Phthalate	10
B75	2,4-Dimethylphenol	10

Analytical Code	Compound	MDL (ppb
733	Semivolatile Organic Analysis (c	ontd)
B76	Dimethyl Phthalate	10
B78	4,6-Dinitro-O-Cresol	10
B80	2,4-Dinitrotoluene	10
B81	2,6-Dinitrotoluene	10
B82	Di-N-Octyl Phthalate	10
B84	1,2-Diphenylhydrazine	10
B88	Fluoranthene	10
B89	Hexachlorobenzene	10
B90	Hexachlorobutadiene	10
B91	Hexachlorocyclopentadiene	10
	Hexachloroethane	10
B92		
B93	Indeno(1,2,3-CD)Pyrene	10
C54	Hexachlorophene	10
C55	Naphthalene	10
C12	Nitrobenzene	10
C17	N-Nitrosodimethylamine	10
C57	Phenol	10
C43	1,2,4-Trichlorobenzene	10
B79	2,4-Dinitrophenol	50
C13	4-Nitrophenol	50
C11	Para-Nitroaniline	50
C28	Pentachlorophenol	50
C44	2,4,5-Trichlorophenol	50
C45	2,4,6-Trichlorophenol	10
C56	1,2,3-Trichlorobenzene	10
C58	1,3,5-Trichlorobenzene	10
C37	1,2,4,5-Tetrachlorobenzene	10
C59	1,2,3,4-Tetrachlorobenzene	10
		10
C60	1,2,3,5-Tetrachlorobenzene	
C26	Pentachlorobenzene	10
B51	Cresols	10
B85	N-Nitrosodinpropylamine	10
C49	Benzo(A)Pyrene	10
C51	Bis(2-Chloroisopropyl) Ether	10
121	Tibutylphosphate	10
B20	Acetophenone	10
B21	Warfarin	10
B22	2-Acetylaminofluorene	10
B23	4-Aminobyphenyl	10
B24	5-(Aminomethyl)-3-Isoxazolol	10
B25	Amitrole	10
B27	Aramite	10
B28	Auramine	10
B29	Benz(C)Acridine	10
B31	Benzene, Dichloromethyl	10
B32	Benzemethoil	. 10

Analytical Code	Compound	MDL (ppb)
733	Semivolatile Organic Analysis (co	ontd)
B36	P-Benzoquinone	10
B37	Benzyl Chloride	10
B43	2-Sec-Butyl-4,6-Dinitrophenol	10
B44	Chloroalkyl Ethers	10
B47	1-Chloro-2,3-Epoxypropane	10
B52	2-Cyclohexyl-4,6-Dinitrophenol	10
B53	Dibenz(A,H)Acridine	10
B54	Dibenz(A,J)Acridine	10
B56	7H-Dibenzo(C,G)Carbazole	10
B57	Dibenzo(A, E)Pyrene	10
B58	Dibenzo(A,H)Pyrene	10
B59	Dibenzo(A, I)Pyrene	10
B66	2,6-Dichlorophenol	10
B68	Dihydrosafrole	10
B69	3,3'-Dimethoxybenzidine	10
B70	P-Dimethylaminoazobenzene	10
B71	7,12-Dimethylbenz(A)Anthracene	10
B72	3,3'-Dimethylbenzidine	10
	Thiofanox	10
B73	Alpha, Alpha-Dimethylphenethyl-	
B74		10
077	Amine	10
B77	Dinitrobenzene	10
B83	Diphenylamine	10
B86	Ethyleneimine	10
B87	Ethyl Methanesulfonate	10
B94	Isosafrole	10
B95	Malononitrile	10
B96	Malphalan	
B97	Methapyrilene	10
B98	Matholonyl	10
B99	2-Methylaziridine	10
C01	3-Methylcholanthrene	10
C02	4,4'-Methylenebis(2-Chloro-	10
	Aniline)	10
C03	2-Methyllactonitrile	10
C05	Methyl Methanesulfonate	10
C06	<pre>2-Methyl-2-(Methylthio)Propion- Aldehyde-0-(Methylcarbonyl)</pre>	
	Oxime	10
C07	Methylthiouracil	10
C07	1,4-Naphthoquinone	10
C08	1-Naphthylamine	10
C09		10
C10	2-Naphthylamine N-Nitrosodi-N-Butylamine	10
C14	N-Nitrosodi-N-Butylamine N-Nitrosodiethanolamine	10
C15		10
C16	N-Nitrosodiethylamine	10

Analytical Code	Compound	OL (ppb)	
733	Semivolatile Organic Analysis (conto	1)	
C18	N-Nitrosomethylethylamine	10	
C19	N-Nitroso-N-Methylurethane	10	
C20	N-Nitrosomethylvinylamine	10	
C21	N-Nitrosomorpholine	10	
22	N-Nitrosonornicotine	10	
3	N-Nitrosonopiperidine	10	
1	Nitrosopyrrolidine	10	
	5-Nitro-O-Toluidine	10	
7	Pentachloronitrobenzene	10	
	Phenacetin	10	
9	Phenylenediamine	10	
	Phthalic Acid Esters	10	
	2-Picoline	10	
	Pronamide	10	
3 4	Reserpine	1.0	
	Rescorzinol	10	
5	Safrol	10	
	2,3,4,6-Tetrachlorophenol	10	
	Thiuram	10	
	Toluenediamine	10	
	O-Toluidine Hydrochloride	10	
	0,0,0-Triethyl Phosphorothicate	10	
	Sym-Trinitrobenzene	10	
	Tris(2,3-Dibromopropyl)-Phosphate	10	
-	Chloronaphazine	10	
	Hexachloropropene	10	
	Kerosene	10	
	Maleic Hydrazide	500	
	Nicotinic Acid	100	
	Strychnine	50	
	Benzo(J)Fluoranthene	10	
	Phosphorous Pesticides		
1	Tetraethylpyrophosphate	2	
	Carbophenothion	2	
	Disulfoton	2	
	Dimethoate	2	
	Methylparathion	2 2 2	
	Ethyl Parathion	2	
	IC Report	500	
72	Nitrate	500	
3	Sulphate	5000	
4	Fluoride	500	
	Chloride	5000	
	Phosphate	1000	

TABLE A.2. (contd)

Analytical Code	Compound	MDL (ppb)
736	Direct Aqueous Injection Analys	is
C53	Hydrazine	3000
C90	Paraldehyde	2000
C94	Acrylamide	10,000
C95	Allyl Alcohol	2500
C97	Chloroacetaldehyde	16,000
C98	3-Chloropropionitrile	4000
H03	Ethyl Carbamate	5000
H04	Ethyl Cyanide	2000)
H09	Isobutyl Alcohol	1000
H11	N-Propylamine	10,000
H12	2-Propyn-1-OL	8000
737	Enhanced Herbicides	
H13	2,4-D	2
H14	2(2,4,5-T)P	2 2 2
H15	(2,4,5-T)	2
39	PCB Analysis	
A54	Arochlor 1016	4.0
A55	Arochlor 1221	1.0
A56	Arochlor 1232	1.0
A57	Arochlor 1242	1.0
A58	Arochlor 1248	1.0
A59	Arochlor 1254	1.0
A60	Arochlor 1260	1.0
20	Arsenic	5.0
21	Mercury	0.1
122	Selenium	5.0
23	Thallium	5.0
\51	Lead	5.0
69	TOC (Total Organic Carbon)	
C70	Cyanide	10.0
C77	Perchlorate	1000
C78	Sulfide	1000
C80	Ammonium Ion	50
C81	Ethylene Glycol	10 ppm
C87	Citrus Red	1000
H16	TC (Total Carbon)	2000
H42	TOX (Total Organic Halogen)	10

TABLE A.3. Sample Container and Preservation Requirements

Constituent(1)	Code	Container Type	Volume (mLs)	Preservative
ICP Metals, Enh(b)	728	Plastic, White Cap	1666	HNOS
Arsenic	A26	Plastic, White Cap	1666	HN03
Selenium	A22	Plastic, White Cap	1666	HNO3
Thallium	A23	Plastic, White Cap	1000	HNO3
Leed, by GFAA	A51	Plastic, White Cap	1666	HNO3
Mercury	A21	Glass, Clear	586	HNO3
Cysnide	C76	Plastic, White Cap	1566	NaOH
Sulfide	C78	Plastic, White Cap	566	ZnAcet/NaOH
Ammonium Ion	C86	Glass, Clear	500	H2S04
TOC	C89	Glass, Amber	258	H3P04
TOX	H42	Glass, Amber Septum	258	H2S04
Pesticides, Enh	729	Glass, Amber	4866	
Herbicides	737	Glass, Amber	4566	
Phosphorous Pasticides	734	Glass, Amber	4866	
4/B/N, Enh	733	Glass, Amber	4666	••
Thioures, Enh	727	Glass, Amber	258	••
Direct Aqueous Inj.	736	Glass, Amber	258	••
Ethylene Glycol	C81	Glass, Amber	256	
Citrus Red	C87	Glass, Amber	256	
TC	H16	Glass, Amber	258	
IC	735	Plastic, White Cap	125	••
Perchlorate	C77	Plastic, White Cap	125	
VOA, Enh	731	Glass, Amber Septum	48	

⁽a) Several apecific constituents reported for some codes.
(b) Enh = enhanced analysis.

APPENDIX B

SAMPLE RESULTS

TABLE B.1. Actual Spring Sample Locations and Analyses Performed - 1988

Area	HRM(a)	Spring No.	River	Analysis
Above 166-B	3.3	3-2		Alpha, Beta, 3H, 735(b)
186-N (N8T)	8.9	NST	x	Alpha, Beta, 3H, 96Sr, Gamma, 9965(b)
Below 166-N	9.5-10.0	••		Aipha, Beta, 3H, 96Sr, Gassa, 735
166-D	11.6	11-1		Alpha, Beta, 3H, 98Sr, 735
100-H	15.0	15-6	X	Alpha, Beta, 3H, U-iso, 9985
Hanford Ferry Landing	25.76	25-4		Alpha, Beta, 3H, 735
Hanford Townsite	27.25	27-2		Alpha, Beta, 3H, 735
Hanford Townsite	27.5	27-3	X	Alpha, Beta, 3H, 98Sr, 99Tc, 735
Hanford Townsite	28.1	28-2	x	Alpha, Bets, 3H, 98Sr, 99Tc, 9985
Hanford Townsite	28.5	28-4		Alpha, Beta, 3H, 98Sr, 99Tc, 735
Wooded Island	38.25	38-1		Alpha, Beta, 3H, 735
Wooded Island	38.8	38-10/39-0		Alpha, Beta, 3H, 735
Above 366	41.5-41.8	41-1/41-2		Alpha, beta, 3H, 735
386	42.0	42-1		Alpha, Beta, 3H, U-iso, 735
386	42.1	42-2	X	Alpha, Beta, SH, U-iso, 236U,, 9985
366	42.3	42-4		Alpha, Beta, 3H, U-iso, 238U, 9985
Port of Benton	43.8	47-1		Alpha, Beta, 3H, 735
Byers Lndg: Irr. Return	(c)	ra(d)		Alpha, Beta, 96Sr, U-iso, 9965
Slide Seep	••	NA		Alpha, Beta, 3H, 98Sr, U-iso, 9985
Ringold Irr. Return		NA		Alpha, Beta, 98Sr, U-iso, 9985
Priest Rapids Dam		NA	x	Alpha, Beta, 3H, 96Sr, 99Tc, U-iso, 9985
Richland Pumphouse		KA	x	Alpha, Beta, 3H, 98Sr, 99Tc, U-iso, 9985

⁽a) HRM = Hanford River Mile.

⁽b) Analytical laboratory code.

⁽c) No data.

⁽d) NA = Not applicable.

TABLE B.2. Columbia River Flow Rates During 1988 Spring Sampling Activities

Date	Flow Rate, cfs
09-04-89	76,000
09-05-89	89,700
09-06-89	96,200
09-07-89	121,000
09-10-89	103,000
09-11-89	46,300
09-12-89	70,800
09-13-89	108,000
09-14-89	99,800
09-15-89	84,700
09-16-89	96,400
09-24-89	102,000
09-25-89	65,200
09-26-89	82,400
09-27-89	92,800
11-12-89	96,800
11-13-89	95,900
11-14-89	100,000
11-15-89	105,000

Background Radionuclide Concentrations Measured in Columbia TABLE B.3 River at Priest Rapids Dam in 1988

	No. of	Concentration (pCi/L)(a)						Drinking Water
Radionuclide(b)	Samples	Max	imum	Min	imum		rage	Standard(c)
Composite System								
Gross Alpha	12	0.85	+ 0.81	-0.07	+ 0.20	0.31	+ 0.17	15
Gross Beta	12	2.31	+ 1.00 .	0.06	± 1.00 + 4	0.96	+ 0.40	50
Tritium	12	89	+ 6	56	+ 4	70	+ 6	20,000
895r	12	0.184	+ 0.084	-0.044	+ 0.072	0.019	+ 0.038	20
905r	12	0.15	+ 0.03	0.05	+ 0.03	0.10	+ 0.02	8
234 _U	12	0.27	+ 0.06	0.11	+ 0.03	0.20	+ 0.03	(d)
23511	12	0.014	+ 0.013	-0.003	+ 0.008	0.006	+ 0.003	
238ŭ	12	0.21	+ 0.04	0.11	+ 0.03	0.17	+ 0.02	
U-Total	12	0.48	\pm 0.07	0.23	± 0.05	0.37	± 0.04	
Continuous System								
60Co P	20	0.0018	+ 0.0019	-0.0012	+ 0.0024	0.0006	+ 0.0003	100
D	20	0.0042	+ 0.0041	-0.0027	+ 0.0042	0.0009	+ 0.0011	
129 _I D	4	0.000045		0.000006		0.000017		1
131 _I p	11	0.0026	+ 0.0037	-0.0011	+ 0.0043	0.0008	+ 0.0008	3
D	11	0.0038	+ 0.0073	-0.0068	+ 0.0114	0.0007	+ 0.0023	
137Cs P	20	0.004	+ 0.0024	0.0002	+ 0.0014	0.0018	+ 0.0005	200
D	20	0.0067	+ 0.0040	-0.0019	+ 0.0044	0.0028	+ 0.0011	
239,240pu p	4	0.00010	+ 0.00008	0.000002		0.00006	+ 0.00004	***
D	4	0.00010	+ 0.00016	0.00002	+ 0.00005	0.00006	+ 0.00004	

(a) Maximum and minimum values ±2 sigma counting error. Average ±2 standard error of the calculated mean.
 (b) Radionuclides measured using the continuous system show the particulate (P) and dissolved (D) fractions separately. Other radionuclides are based on samples collected by the composite system.
 (c) From State of Washington and EPA.

(d) Dashes indicate no concentration guides provided in DWS.

Columbia River Background Water Quality Data for 1988 (from Jaquish and Bryce 1989) TABLE B.4

			Vernita B	Bridge (Ups	tream)
Analysis	Units	No. of Samples	Maximum	Minimum	Annual (a)
PNL Environmental Monitoring					
рН	pH units	12	8.5	7.4	NA(b)
Fecal coliform	#/100 mL		130	2	2(c)
Total coliform	#/100 mL	12	1600	2 2	48 (c)
Biological oxygen demand	mg/L	12	5.2	0.7	2.1 + 0.8
Nitrate	mg/L	12	0.23	0.05	0.14 ± 0.03
USGS Sampling Program(d)					
Temperature(e)	°C	365	19.6	1.8	11.3
Dissolved oxygen	mg/L	6	13.4	8.8	11.5 + 1.4
Turbidity	NTU	6	1.8	0.4	1.0 + 0.4
pH	pH units	6 .	8.8	8.0	NĀ 0.4
Fecal coliform	#/100mL	6	3	<1	2(c)
Suspended solids, 105°C	mg/L	NR		•	21-7
Dissolved solids, 180°C	mg/L	6	88	71	81 + 6
Specific conductance	umhos	6	162	123	81 + 6 140 + 15 68 + 7
Hardness, as CaCO3	mg/L	6	77	58	68 + 7
Phosphorus, total	mg/L	6	0.03	0.02	0.023 ± 0.004
Chromium, dissolved	µg/L	6 3	<1	<1	<1
Nitrogen, Kjeldahl	mg/L		0.5	<0.2	<0.28 + 0.11
Total organic carbon	mg/L	6 4	2.8	1.4	2.1 + 0.7
Iron, dissolved	µg/L	3 5	65	9	$\frac{28}{4}$
Ammonia, dissolved (as N)	mg/L	5	0.05	<0.01	<0.02 ∓ 0.02

⁽a) Average values +2 standard error of the calculated mcan.
(b) NA = Not applicable.
(c) Annual median.
(d) Provisional data subject to revision.
(e) Maximum and minimum represent daily averages.

TABLE B.5. Estimated Background Levels for Selected Constituents in Hanford Ground Water (from Evans et al. 1989)

Constituent	Detection Limit(a)	Background Concentration(a)
Aluminum	2(b)	(b)
Ammonia	50	<50
Arsenic	0.2(b)	3.9 + 2.4(b)
Barium	* 6	42 + 20
Beryllium	0.3(b)	<0.3(b)
Bismuth	0.02(b)	<0.02(b)
Boron	50(b)	<50(b)
Cadmium	0.2(b)	<0.2(b)
Calcium	50	$40,000 \pm 10,300$
Chloride	500	$10,300 \pm 6,500$
Chromium	2(b)	$4.0 \pm 2.0(b)$
Copper	1(b)	<1(b)
Cyanide	10	<10
Fluoride	500	370 + 100
Lead	0.5(b)	<0.5(b)
Magnesium	10	11,800 + 3,400
Manganese	5	7 <u>+</u> 5
Mercury	0.1	<0.1
Nickel	4(b)	<4(b)
Phosphate	1,000	<1,000
Potassium	100	4,950 + 1,240
Selenium	2(b)	<2(b)
Silver	10	<10
Sodium	10	18,260 + 10,150
Strontium	20	236 ± 102
Sulfate	500	34,300 ± 16,900
Jranium	0.5(c)	$1.7 \pm 0.8(c)$
/anadium	5	17 <u>+</u> 9
linc	5	6 + 2

TABLE B.5. (contd)

Constituent	Detection Limit(a)	Background Concentration(a)		
Alkaïinity		123,000 ± 21,000		
pH		7.64 ± 0.16		
Total Organic Carbon	200	586 ± 347		
Conductivity	1(d)	$380 \pm 82(d)$		
Gross Alpha	0.5(c)	2.5 ± 1.4(c)		
Gross Beta	4(c)	$19 \pm 12(c)$		
Radium	0.2(c)	<0.2(c)		

Units in ppb unless otherwise noted. Based on ICP/MS data. Units in pCi/L data. Units in µmho/cm.

TABLE B.6. Radiological Analyses of Water Samples Collected from the Columbia River, Columbia River Shoreline Springs (onsite) and Irrigation Wastewater Discharges (offsite)

(All Concentrations in pCi/l. Blank spaces indicate not analyzed.)

Sample	Sasple	Sample				
Date	Location	Source	Gross Alpha	Gross Beta	Tritium	
11/14/88	HRM 2.5	Spring	Ø.49 +/- Ø.38	1.37 +/- 1.67	-320 -/-	128
11/14/88	HRM 3.3	Spring	1.67 -/- 6.96	7.87 +/- 1.93	1156 -/-	162
69/15/88	HRM 8.9	River			76496 +/-	916
69/15/88	HRM 8.9	Spring	0.00035 -/- 0.28	16866 -/- 296	75886 -/-	968
69/66/88	HRM 8.9	Spring	0.55 +/- 0.61	13866 -/- 186	74888 +/-	700
69/66/88	HRM 9.5	Spring	8.37 +/- 8.48	73.8 +/- 7.8	111606 -/-	868
11/14/88	HRM 11.6	Spring	6.30 +/- 6.33	1.65 +/- 1.12	21.5 -/-	137
69/12/88	HRM 15.6	River			65.7 -/-	127
69/12/88	HRM 15.6	Spring	6.35 -/- 6.35	1.63 -/- 1.27	545 -/-	178
69/12/88	HRM 25.75	Spring	6.42 -/- 6.42	6.69 -/- 1.14	264 -/-	169
69/12/88	HRM 27.25	Spring	2.50 -/- 2.67	4.33 -/- 4.1	7428 -/-	296
69/12/88	HRM 27.5	River			26466 -/-	525
69/12/88	HRM 27.5	Spring	2.11 +/- 1.02	14.2 +/- 2.71	72668 •/-	888
69/12/88	HRM 28.1	River			158666 -/- 1	256
69/12/88	HRM 28.1	Spring	2.32 -/- 1.67	48 +/- 4.88	155666 -/- 1	290
69/66/68	HRM 28.1	Spring	2.62 -/- 1.12	168 -/- 11	143666 •/-	986
69/12/88	HRM 28.5	Spring	1.49 -/- 6.91	45 -/- 4.65	145666 -/- 1	256
69/26/88	HRM 38.25	Spring	2.91 -/- 1.14	1.85 +/- 1.4	2636 -/-	231
69/26/88	HRM 38.8	Spring	2.28 -/- 1.11	6.84 +/- 1.98	382 -/-	182
69/26/88	HRM 41.58	Spring	3.25 +/- 1.49	16.1 -/- 2.41	6586 -/-	388
69/26/88	HRM 42.6	Spring	4.42 +/- 1.18	5.25 +/- 1.69	1676 -/-	192
69/12/88	HRM 42.1	River			485 +/-	176
69/16/88	HRM 42.1	Spring	7.95 +/- 1.46	11.6 +/- 3.3	168 -/-	115
69/12/88	HRM 42.1	Spring	8.48 +/- 1.29	7.31 -/- 1.94	348 -/-	172
69/12/88	HRM 42.3	Spring	8.51 +/- 1.38	9.61 +/- 2.22	283 -/-	176
69/26/68	HRM 43.8	Spring	6.52 +/- 8.66	4.69 +/- 1.78	64.8 -/-	163
69/15/88	Byers Lndg	Irr. Rtn	8.75 ·/- 8.46	6.39 +/- 2.12		
69/15/88	Slide	Seep	2.62 -/- 6.99	2.68 +/- 1.88	377 -/-	174
69/15/88	Ringold	Irr. Rtn	1.76 -/- 6.73	9.94 -/- 2.51		
69/13/88	Priest Rapids	River			186 -/-	167
99/13/88	Richland	Pumphouse			581 -/-	179

TABLE B.6. (contd)

Sample	Sample	Sample			
Date	Location	Source	Sr96	Co66	Zn65
11/14/88	HRM 2.5	Spring		-0.69 +/- 3.39	2.63 -/- 9.41
11/14/88	HRM 3.3	Spring		0.42 +/- 1.71	-1.25 +/- 5.14
69/15/88	HRM 8.9	River	6746 +/- 186	38.9 +/- 4.53	3.25 +/- 6.72
69/15/88	HRM 8.9	Spring	7270 -/- 192	53.4 +/- 19.4	5.30 -/-15.60
69/66/88	HRM 8.9	Spring	6688 +/- 269	45 +/- 4.5	6.76 -/- 2.98
69/66/88	HRM 9.5	Spring	6.31 -/- 6.89	36.3 +/- 4	-2.16 -/- 3.36
11/14/88	HRM 11.0	Spring	1.41 -/- 6.43	-2.66 +/- 3.46	-1.53 -/- 9.94
59/12/88	HRM 15.8	River			
69/12/88	HRM 15.6	Spring		-6.55 +/- 1.92	-0.38 +/- 8.25
69/12/88	HRM 25.75	Spring		-\$.682 ·/- 1.18	-1.38 -/- 6.41
69/12/88	HRM 27.25	Spring			
69/12/88	HRM 27.5	River			
69/12/88	HRM 27.5	Spring	-\$.6693 •/- \$.33	1.67 -/- 3.88	-4.41 +/- 16.1
69/12/88	HRM 28.1	River			
69/12/88	HRM 28.1	Spring	8.674 +/- 8.35	4.72 +/- 4.76	1.52 -/- 14.4
69/96/88	HRM 28.1	Spring	8.79 ·/- 8.11	4 +/- 1.8	-8.20 ·/- 2.60
69/12/88	HRM 28.5	Spring	€.6614 +/- €.33	2.82 -/- 3.16	4.39 -/- 8.77
69/26/88	HRM 38.25	Spring		-1.66 -/- 2.63	-4.32 -/- 5.49
69/26/88	HRM 38.8	Spring		-6.71 -/- 1.8	-1.77 +/- 6.13
69/28/88	HRM 41.58	Spring		0.38 +/- 1.45	-2.14 -/- 5.79
69/26/86	HRM 42.5	Spring		€.34 •/- 1.53	4.63 -/- 5.12
69/12/88	HRM 42.1	River		6.61 ·/- 1.42	1.68 -/- 5.86
69/16/88	HRM 42.1	Spring	5.16 */- 6.67	€.25 •/- €.3	-6.95 +/- 1.66
69/12/88	HRM 42.1	Spring		-3.38 +/- 2.74	5.44 -/- 7.98
69/12/88	HRM 42.3	Spring		1.61 -/- 2.18	2.96 +/- 7.25
69/26/88	HRM 43.6	Spring		8.41 +/- 1.29	-2.21 -/- 8.65
69/15/88	Byers Lndg	Irr. Rtn	0.422 ·/- 0.35	1.26 +/- 2.48	8.69 +/− 7.84
#9/15/88	Slide	Seep	■.8897 •/- ■.35	2.95 +/- 3.81	3.35 +/- 11.4
69/15/88	Ringold	Irr. Rtn	€.385 •/- €.38	1.57 +/- 3.51	-4.48 +/- 8.71
69/13/86	Priest Rapids	River	-6.172 •/- 6.29	-2.35 -/- 2.88	-4.44 +/- 7.73
69/13/88	Richland	Pumphouse	6.28 ·/- 6.35	-8.49 -/- 4.67	-2.54 +/- 14.3

TABLE B.6. (contd)

Sample	Sample	Sample			
Date	Location	Source	Tc99	Ru186	Sb125
11/14/88	HRM 2.5	Spring		12.4 +/- 31.8	
11/14/88	HRM 3.3	Spring		9.67 +/- 15.6	
69/15/88	HRM 8.9	River		11.8 -/- 24.2	28.6 +/- 7.51
69/15/88	HRM 8.9	Spring		12.6 -/- 71.2	
69/66/88	HRM 8.9	Spring		8.9 +/- 11.4	43.1 +/- 6.4
69/66/88	HRM 9.5	Spring		13.5 +/- 14.8	
11/14/88	HRM 11.0	Spring		-4.62 -/- 33.7	
69/12/88	HRM 15.6	River			
69/12/88	HRM 15.6	Spring		-1.69 -/- 26.1	
69/12/88	HRM 25.75	Spring		6.88 +/- 15.8	
69/12/88	HRM 27.25	Spring			
69/12/88	HRM 27.5	River			
09/12/88	HRM 27.5	Spring	48.4 -/- 1.76	3.22 +/- 39	
69/12/88	HRM 28.1	River			
69/12/88	HRM 28.1	Spring	223 +/- 2.95	-2.65 +/- 41.4	
69/66/88	HRM 28.1	Spring	228 -/- 3	3.8 -/- 7.6	
69/12/88	HRM 28.5	Spring	215 +/- 2.89	24.1 +/- 28.7	
69/25/88	HRM 38.25	Spring		-2.62 +/- 17.1	
69/25/88	HRM 38.6	Spring		8.93 +/- 26.6	
65/25/88	HRM 41.58	Spring		-3.9 -/- 21.8	
69/20/88	HRM 42.6	Spring		13.2 -/- 17.6	
69/12/63	HRM 42.1	River		9.6 +/- 17.4	
69/16/88	HRM 42.1	Spring		-6.3 -/- 6.1	
69/12/88	HRM 42.1	Spring		9.29 +/- 21.7	
69/12/88	HRM 42.3	Spring		-16.8 -/- 27.1	
69/26/88	HR4 43.8	Spring		11.3 +/- 19.9	
69/15/88	Byers !.ndg	Irr. Rtn		2.37 -/- 24.6	
69/15/88	Slide	Seep		12.6 +/- 38.7	
69/15/88	Ringold	Irr. Rtn		-4.72 +/- 33.8	
69/13/88	Priest Rapids	River	6.63 +/- 1.16	-14.8 +/- 26.8	
69/13/88	Richland	Funphouse	1.77 +/- 1.24	41.8 +/- 48.4	

TABLE B.6. (contd)

	Sample	Sample	Sample			
	Date	Location	Source	Ca137	U234	U235
ľ	11/14/88	HRM 2.5	Spring	1.16 +/- 2.7		
	11/14/88	HRM 3.3	Spring	-0.62 +/- 1.37		
	69/15/88	HRM 8.9	River	-6.66 +/- 1.63		
	69/15/88	HRM 8.9	Spring	-6.37 -/- 4.35		
	69/66/88	HRM 8.9	Spring	-8.2 -/- 1		
	69/65/88	HRM 9.5	Spring	0.41 +/- 1.67		
	11/14/88	HRM 11.6	Spring	-0.19 +/- 3.5		
	69/12/88	HRM 15.5	River			4
	69/12/88	HRM 15.0	Spring	-6.94 -/- 1.62	0.083 +/- 0.03	-6.8631 -/- 0.8841
	69/12/88	HRM 25.75	Spring	8.41 •/- 1.43		
	69/12/88	HRM 27.25	Spring			
	69/12/88	HRM 27.5	River			
	69/12/88	HRM 27.5	Spring	€.63 +/- 2.78		
	69/12/88	HRM 28.1	River			
	69/12/88	HRM 28.1	Spring	-3.74 +/- 3.31		
	69/66/88	HRM 28.1	Spring	-8.5 +/- 1.1		
	69/12/88	HRM 28.5	Spring	-1.3 -/- 2.58		
	69/26/68	HRM 38.25	Spring	-0.58 -/- 1.41		
	69/26/88	HRM 38.8	Spring	1.37 +/- 1.79		
	69/26/88	HRM 41.58	Spring	0.45 -/- 1.76		
	69/26/88	HRM 42.6	Spring	-1.64 +/- 1.53	2.63 -/- 6.13	6.18 -/- 6.641
	69/12/88	HRM 42.1	River	-6.94 +/- 1.25	4.28 -/- 0.21	0.31 -/- 0.858
	69/16/88	HRM 42.1	Spring	6.4 ·/- 6.6	4.96 -/- 6.28	8.26 ·/- 8.64
	69/12/88	HRM 42.1	Spring	-6.44 ·/- 1.99	4.48 -/- 6.20	6.38 +/- 8.858
	69/12/88	HRM 42.3	Spring	-8.36 -/- 2.66	3.48 -/- 6.18	6.24 -/- 6.548
	09/26/88	HRM 43.6	Spring	6.49 -/- 1.65		
	09/15/88	Byers Lndg	Irr. Rtn	6.76 -/- 2.83	2.6 -/- 5.25	0.15 -/- 0.049
	69/15/88	Slide	Seep	2.51 -/- 3.02	4.8 -/- 6.26	0.14 -/- 0.634
	69/15/88	Ringeld	Irr. Rtn	6.88 -/- 2.99	3.62 -/- 6.16	8.17 -/- 8.648
	69/13/88	Priest Rapids	River	-1.61 -/- 2.5	8.16 +/- 8.84	0.0082 -/- 0.013
	69/13/88	Richland	Pumphouse	-6.27 +/- 3.91	0.18 +/- 0.04	-0.0007 +/- 8.889

TABLE B.6. (contd)

Sample	Sample	Sample	
Date	Location	Source	U238
11/14/88	HRM 2.5	Spring	
11/14/88	HRM 3.3	Spring	
69/15/88	HRM 8.9	River	
69/15/88	HRM 8.9	Spring	
69/55/88	HRM 8.9	Spring	
59/56/88	HRM 9.5	Spring	
11/14/88	HRM 11.0	Spring	
69/12/88	HRM 15.0	River	
69/12/88	HRM 15.6	Spring	0.097 +/- 0.036
69/12/88	HRM 25.75	Spring	
69/12/88	HRM 27.25	Spring	
69/12/88	HRM 27.5	River	
69/12/88	HRM 27.5	Spring	
69/12/88	HRM 28.1	River	
69/12/88	HRM 28.1	Spring	
99/56/88	HRM 28.1	Spring	
69/12/88	HRM 28.5	Spring	
69/28/88	HRM 38.25	Spring	
69/26/88	HRM 38.8	Spring	
69/26/88	HRM 41.68	Spring	
69/26/88	HRM 42.6	Spring	1.94 -/- 5.13
69/12/88	HRM 42.1	River	3.95 +/- 0.20
69/16/88	HRM 42.1	Spring	4.48 •/- 6.26
59/12/88	HRM 42.1	Spring	4.8 -/- 6.25
69/12/88	HRM 42.3	Spring	3 -/- 0.17
59/26/88	HRM 43.6	Spring	
69/15/88	Byers Lndg	Irr. Rtn	2.27 -/- 6.19
69/15/88	Slide	Seep	3.91 •/- 6.18
69/15/88	Ringold	Irr. Rtn	2.32 -/- 6.14
69/13/88	Priest Rapids	River	6.13 -/- 6.836
99/13/88	Richland	Pumphouse	5.16 ·/- 5.64

TABLE B.7. Nonradiological Analysis of Water Sample Collected from the Columbia River, Columbia River Shoreline Springs (onsite) and Irrigation Water Discharges (offsite)

(All Concentrations in ppb. Blank spaces indicate not analyzed.)

		Detection	88316	88322	88335	88346	88348
Analytical		Limit	RM 8.9	RM 14.5	RM 28.1	RM 42.1	RM 42.3
Code	Compound	(ppb)	Spring	Spring	Spring	Spring	Spring
726	ICP Metals						
AØ1	beryllium	5	<5	⟨5	<5	<5	<5
AS3	Strontium	20	156	132	333	169	119
A84	Zinc	5	47	11	17	23	16
ASS	Calcium	56	41797	23986	45526	24289	26492
ASS .	Berium	6	52	43	66	48	54
A67	Cadaius	2	<2	<2	(2	<2	(2
ASS.	Chronium	16	<16	<10	(16	<16	<16
A16	Silver	16	(16	<16	<16	(10	<16
A11	Sodium	266	4337	3165	21435	15269	12320
A12	Nickel	16	<10	(16	<10	<10	(10
A13	Copper	10	<16	<10	<16	34	<10
A14	Vanadium	5	< 5	<6	16	< 5	< 5
A15	Antieony	100	<166	<100	<100	<165	<100
A16	Aluminum	156	(158	<158	366	<159	<15€
A17	Manganese	5	6	7	24	< 5	5
A18	Potassius	100	1663	923	2784	2493	2461
A19	Iron	36	163	⟨3€	451	87	121
ASS	Magnes i un	50	6614	5467	13127	4584	4821
727	Enhanced Thioures						
A24	Thioures	266	<200	(200	<286	<200	⟨299
A25	1-Acety I-2-Thioures	266	<200	(200	(200	<266	<29€
A26	1-(S-Chiorophenyi)-Thioures	266	(286	(200	(286	<296	<20€
A27	Diethylstilbestrol	299	<200	(200	(200	<286	<296
A28	Ethy leneth ioures	286	<266	(200	(200	<266	<255
A29	1-Naphthy I-2-Thioures	266	<266	<200	(200	<29€	⟨286
A32	N-Pheny Ithiourea	500	(586	(588	<586	<500	<500
729	Enhanced Pesticides						
A33	Endrin	6.1	< 6.1	<0.1	<0.1	<0.1	⟨6.1
A34	Methoxychlor	3	<3	⟨3	⟨3	⟨3	<3
A35	Toxaphene	1	<1	<1	<1	<1	<1
A36	Alpha BHC	6.1	⟨∅.1	<0.1	< 6.1	⟨∅.1	⟨∅.1
A37	Beta BHC	6.1	<0.1	< 6.1	< 6.1	< 6.1	⟨8.1
A38	Gamma BHC	6.1	⟨∅.1	⟨∅.1	<0.1	<0.1	<8.1
A39	Delta BHC	6.1	<0.1	<0.1	<0.1	<0.1	<0.1
A45	DDD	●.1	<0.1	<0.1	⟨∅.1	< 6.1	<0.1
A41	DDE	6.1	< 6.1	<0.1	<0.1	<0.1	<0.1
A42	DDT	6.1	< 6.1	<0.1	<0.1	<9.1	<0.1
A43	Heptachlor	0.1	< 6.1	<6.1	<0.1	<8.1	⟨∅.1
A44	Heptachlor Epoxide	6.1	<0.1	⟨∅.1	⟨∅.1	<0.1	<0.1

TABLE B.7. (contd)

Analytical Code	Compound	Detection Limit (ppb)	88351 Byers Lndg Irr Rtn	88353 Slide Seep	88355 Ringold Irr Rtn	88357 Priest Rapid River	88359 Richland Pumphouse	
726	ICP Metals							
AØ1	Beryllium	5	< 5	< 5	(5	₹5	(5	
A63	Strontium	20	323	1045	379	125	126	
AB4	Zinc	5	8	10	6	12	9	
AØ5	Calcium	50	45680	93019	52916	21657	21362	
AØ6	Barius	6	83	64	53	33	32	
A67	Cadeium	2	<2	<2	<2	<2	<2	
ABA	Chromium	10	<10	<10	(16	<16	<10	
A16	Silver	10	<10	<10	(15	<10	<10	
A11	Sodium	200	42975	72148	47725	2452	2563	
A12	Nickel	16	(10	(18	(16	(10	(18	
A13	Copper	16	(16	(10	(10	<16	(19	
A14	Vanadius	5	13	8	15	(5	(5	
A15	Antimony	100	(190	(100	(100	<100	(100	
A16	Aluminum	156	(150	692 93	(15 5	(150 14	<15 0	
A17	Manganese	V. N. J.	16				862	
A18	Potassium	160	5067	2605	7316	811	37	
A19 A5Ø	Iron Magnesium	3 0	155 21418	596 48425	2 00 25446	160 4777	4567	
	4							
727	Enhanced Thioures		1000	4000	1000	1088	/000	
A24	Thioures	200	(200	(200	<200	(200	(200	
A25	1-Acetyl-2-Thiourea	206	<200	<200	<200 <200	<200 <200	(200	
A26	1-(8-Chlorophenyl)-Thioures	200	(200	<200 <200	(200	(200	(200	
A27	Diethylstilbestrol	200	<200 <200	(200	<200	(200	(200	
A28 A29	Ethylenethioures	200	(200	(200	(200	(200	(200	
A32	1-Naphthy I-2-Thiourea N-Pheny Ithiourea	500	(506	(588	<500	<500	<500	
729	Enhanced Pesticides							
A33	Endrin	6.1	(0.1	⟨∅.1	⟨∅.1	⟨∅.1	(0.1	
A34	Methoxychlor	3	(3	(3	(3	(3	(3	
A35	Toxaphene	1	(1	<1	(1	(1	(1	
A36	Alpha BHC	6.1	⟨∅.1	<0.1	⟨∅.1	⟨∅.1	(0.1	
A37	Beta BHC	9.1	⟨€.1	⟨∅.1	⟨∅.1	(0.1	⟨∅.1	
		0.1	⟨∅.1	⟨∅.1	(9.1	(0.1	(0.1	
A38	Gamma BHC Deita BHC	Ø.1	⟨∅.1	⟨€.1	(8.1	(0.1	(6.1	
A39 A48	DOD BHC	8.1	⟨∅.1	(8.1	(8.1	(0.1	(6.1	
					(0.1	(0.1	(0.1	
A41	DDE	0.1	<0.1 <0.1	<0.1 <0.1	⟨∅.1	(0.1	(0.1	
A42 A43	Heptachior	6.1	(0.1	(0.1	(0.1	(0.1	(0.1	
A43	пертасптог	9.1	(0.1	(0.1	(0.1	(0.1	(0.1	

TABLE B.7. (contd)

Analytical Code	Compound	Detection Limit (ppb)	88316 RM 8.9 Spring	88322 RM 14.5 Spring	88335 RM 28.1 Spring	88345 RM 42.1 Spring	88348 RM 42.3 Spring	
A46	Dieldrin	0.1	⟨€.1	⟨€.1	⟨∅.1	⟨€.1	⟨€.1	
M7	Aldrin	0.1	⟨€.1	<0.1	<0.1	<0.1	⟨∅.1	
A48	Chiordane	1	<1	<1	<1	<1	<1	
A49	Endosulfan I	■.1	⟨€.1	⟨€.1	<0.1	<0.1	<0.1	
A52	Endosulfan II	0.1	⟨∅.1	⟨€.1	<0.1	<0.1	<0.1	
C62	Chlorobenzilate	30	(38	⟨3€	⟨3€	⟨35	⟨3€	
X10	DBC	1	<1	(1	< 1	(1	(1	
731	Volatile Organic Compounds						15	
A61	Tetrach loromethane	. 5	(5	(8	(5	(5	(5	
A62	Benzene		(5	₹5	(5	(5	(5 66	
A63	Dioxane	500	(566	<566	(500	(586 (18	(18	
A84	Methylethyl Ketone	10	(10	(10	(18 (586	(588	(500	
A85	Pyridine	580	(588	(500	(5	(5	(5	
ASS	Toluene	8	(5	(5	(5	(5	(5	
A67	1,1,1-Trichloroethane	8	(5	(5	(5	(5	(5	
A68	1,1,2-Trichloroethane	5	(5	(5	(5	(5	(5	
A69	1,1,2-Trichloroethene	5	(5	(5	(5	(5	(5	
A76	Perch loroethy lene	5		(5	(5	(5	(5	
A71	Xylene (0, P)	5	<5	(5	(5	(5	(5	
814	Xylene (M)	5	(5	(10	(10	(16	(10	
A76	Methyl Broside	10	(10	(10	(10	(10	(10	
A77	Carbon Disulfide	16	(10	(10	(10	(15	(10	
A78	Chiorobenzene	10	(15	(18	(10	(10	(10	
A79	2-Chloroethylvinylether	10	<10 <5	(16	(5	24	19	
ABO	Chloroform	5	• •	(10	(10	(10	(10	
A81	Chloromethane	10	<10 <10	(10	(10	(18	(10	
A89	1,1-Dichloroethane	10	(10	(10	(10	(18	(10	
A98	1,2-Dichloroethane	10	(10	(10	(10	(10	(10	
A91	Trans-1,2-Dichloroethene	16	(10	(10	(10	(10	(10	
A92	1,1-Dichlorethene	10	(10	(15	(10	(10	(10	
A93	Methylene Chloride	16	(18	(10	(10	(10	(10	
A94	1,2-Dichloropropane	15	<10	(10	(10	(10	(10	
A95	1,3-Dichloropropenes	10	(10	(10	(10	(10	(10	
B66	1,1,2,2-Tetrachloroethane	10	(10	(10	(10	(10	(10	
968	Bronoform	10	(10	(10	(10	(10	(10	
813	Vinyl Chloride	10	(10	(15	(10	(10	<18	
1168	Hexone	16	<10	. (10	(10	(18	(10	
A72	Acrolein	10	(10	(10	(10	(18	<10	
A73	Acrylonitrile Bis (Chloromethyl) Ether	16	(10	(10	(10	(10	(18	

TABLE B.7. (contd)

Analytical Code	Compound	Detection Limit (ppb)	88351 Byers Lndg Irr Rtn	88353 Slide Seep	88355 Ringold Irr Rtn	88357 Priest Rapid River	88359 Richland Pumphouse
729	Enhanced Pesticides		**********				
A46	Dieldrin	5.1	⟨€.1	⟨€.1	⟨€.1	⟨€.1	⟨€.1
A47	Aldrin	. 6.1	⟨€.1	⟨€.1	⟨€.1	⟨8.1	⟨6.1
A48	Chlordane	1	<1	<1	(1	<1	<1
A49	Endosulfan I	6.1	⟨€.1	⟨€.1	(6.1	<0.1	⟨∅.1
A52	Endosulfan II	6.1	₹8.1	<0.1	⟨∅.1	<0.1	⟨∅.1
C82	Chlorobenzilate	30	⟨3₽	<3€	⟨3€	⟨3€	⟨3€
X16	DBC	1	<1	<1	(1	<1	(1
731	Volatile Organic Compounds						
A61	Tetrachloromethane	5	(5	(5	. <5	<5	<5
A62	Benzene	. 5	⟨5	(5	(5	<5	<5
A63	Dioxane	500	<566	<566	(500	<566	<566
A84	Methylethyl Ketone	16	₹16	<10	(10	<10	<10
A65	Pyridine	566	<566	<586	<566	<500	<500
A86	Toluene	5	<5	(5	<5	<5	(5
A67	1,1,1-Trichloroethane	8	< 5	<5	<5	<5	(5
A86	1,1,2-Trichloroethane	5	< 5	< 5	(5	₹5	< 5
A69	1,1,2-Trichloroethens	5	(5	< 5	(5	(5	(5
A76	Perchloroethylene	5	< 5	<5	<5	<5	<5
A71	Xylene (0, P)	5	₹5	(5	<5	<5	(5
814	Xylene (M)	5	<5	< 5	(5	₹5	<5
A75	Methy: Bromide	10	<10	<16	<18 .	<16	(10
A77	Carbon Disulfide	16	<10	<16	<10	<16	(18
A78	Chiorobenzene	16	<10	<16	<10	<16	(16
A79	2-Chloroethylvinylether	10	<10	<10	<10	<10	<10
A80	Chlorofore	5	<5	<8	<5	<5	<5
A81	Chloromethane	10	<10	<10	<16	<10	<15
A89	1,1-Dichloroethane	10	<10	<10	<10	<16	(10
ASS	1,2-Dichloroethane	10	<10	(16	<10	<10	<10
A91	Trans-1,2-Dichloroethene	10	<16	<10	<10	<16	(16
A92	1,1-Dichlorethene	. 10	<10	. (16	<10	<10	(10
A93	Methylene Chloride	10	<10	<18	<10	<10	<10
A94	1,2-Dichloropropane	10	<10	(16	<10	<16	<10
A95	1,3-Dichloropropenes	10	<10	(10	<10	<10	(18
B#6	1,1,2,2-Tetrachtoroethane	10	<10	(10	<10	<10	<18
BØ8	Brosoform	16	(10	(10	(10	(18	<18
B13	Vinyl Chloride	10	(10	<18	<10	<10	(19
H68	Hexone	10	<10	(10	<10	<10	<18
A72	Acrolein	16	<16	<10	<16	<18	<18
A73	Acrylonitrile	10	<10	(10	<10	<10	<18
A74	Bis (Chloromethyl) Ether	16	<10	<10	(10	<10	<18

TABLE B.7. (contd)

		Detection	88316	88322	88335	88346	88348
Analytical	-	Limit	RM 8.9	RM 14.5	RM 28.1	RM 42.1	RM 42.3
Code	Compound	(ppb)	Spring	Spring	Spring	Spring	Spring
731	Volatile Organic Compounds		7				
A75	Broso Acetone	10	<10	<10	<10	<10	(18
A82	Chioromethy lasthy lether	10	<10	<10	<10	(10	(10
A83	Crotona I dehyde	10	<10	<10	(10	<10	(10
A84	1,2-Dibroso-3-Chloropropane	10	<10	<10	<10	<10	(10
A85	1,2-Dibroscethane	10	<10	<10	<10	<10	(10
A86	Dibrososthans	10	<10	<18	<10	<10	<10
A87	1,4-Dichloro-2-Butene	10	<10	<10	<10	<10	<10
A88	Dichlorodifluoromethane	19	<10	<10	<10	<16	(10
A96	N, N-Diethy lhydrazine	10	<10	(10	<10	<10	(10
A99	Hydrogen Sulfide	10	(10	<18	(10	<10	(10
861	Iodo methane	10	(16	(18	<10	(10	(18
B02	Methacrylonitrile	10	(10	<10	(10	<10	(10
863	Methanethiol	10	(10	<10	(10	(10	(10
B84	Pentach lo roethane	16	(10	<10	<10	<10	(10
B65	1,1,1,2-Tetrachioroethane	10	(10	<10	<10	<16	(10
809	Trichlo. osethanethiol	10	(16	<16	<10	<10	(16
B16	Trichlorof luoromethane	10	(16	<10	<10	<10	(16
B11	Trichloropropene	10	(10	<10	<10	<16	(10
B12	1,2,3-Trichloropropane	16	(18	<10	<18	<18	(10
815	Diethylarsine	10	(10	<10	<10	(10	(10
C71	Forez I dehyde	500	(500	(586	(500	<588	<588
CS4	Methyl Methacrylate	16	(10	<10	<10	<10	(16
H86	Ethyl Methacrylate	10	(10	<10	<10	(10	<10
B19	Acetonitrile	3 ppe	(3 ppm	(3 ppm	<3 ppm	(3 ppm	(3 ppe
HØ5	Ethylene Oxide	3 ppm	(3 ppm	(3 ppm	(3 ppm	(3 ppm	(3 ppe
733	Semi-Volatile Organic Compounds						
826	Aniline	< 10	< 10	< 18	< 18	< 10	< 16
B30	Benz (A) Anthracane	< 10	< 10	< 10	< 10	< 10	< 10
B33	Benzidine	< 10	< 10	(10	< 10	< 10	(10
B34	Benzo (B) Fluoranthene	< 10	< 10	< 10	< 10	< 10	< 10
B38	Bis (2-Chloroethoxy) Methane	< 10	< 10	< 10	< 10	< 10	< 16
839	Bis (2-Chloroethy !) Ether	(10	< 10	< 10	< 10	< 10	< 10
B46	Bis (2-Ethylhexyl) Phthalate	< 10	< 10	< 10	< 10	< 10	< 16
B41	4-Brosopheny! Pheny! Ether	< 16	(16	< 10	< 10	< 10	< 16
B42	Butyl Benzyl Phthalate	< 10	< 10	< 10	< 10	< 10	< 10
B45	P-Chloroaniline	< 10	< 10	< 10	< 10	(10	(10
B46	P-Chloro M-Cresol	< 16	< 10	< 10	< 10	(18	< 10
B48	2-Chioronaphthaiene	< 10	< 16	(16	< 16	(16	< 10
B49	2-Chiorophenol	< 10	< 10	< 10	< 16	< 10	< 10
856	Chrysene	< 16	< 16	< 10	< 16	< 10	(10

TABLE B.7. (contd)

Analytica Code	Compound	Detection Limit (ppb)	88351 Byers Lndg Irr Rtn	88353 Slide Seep	88355 Ringold Irr Rtn	88367 Priest Rapid River	88359 Richland Pumphouse
731	Volatile Organic Compounds		41.6	41.0	410	410	218
A75	Broso Acetone	16	(16	(10	(10	(18	(10
A82	Chloromethy laethy lether	16	(10	(10	(10	(10	(10
A83	Crotonal dehyde	10	(10	<18 <18	<18 <18	(10 (10	(18
A85	1,2-Dibroso-3-Chloropropane 1,2-Dibrosoethane	16	(10	(18	(10	(10	
A86	Dibrososethane	10	(18	<10	(10	(16	(10
A87	1.4-Dichloro-2-Butene	10	(10	(16	(18	(16	(10
A88	Dichlorodifluoromethans	10	(16	<10	(18	(10	(16
A96	N,N-Diethylhydrazine	16	(10	(16	(16	(16	(16
A99	Hydrogen Sulfide	16	(16	(16	(16	(10	(16
861	Iodo methane	16	(16	(16	(16	(16	(16
			•	•		•	Mark Co.
B02	Methacry Ionitri le	10	(16	(16	(16	(16	(16
B63 B64	Methanethici Pentachioroethane	16	(10	(16	(16 (16	(16	<16 <16
B#5		10	•	(16	(16	(16	(16
	1,1,1,2-Tetrachloroethane		(10	•		•	
B09	Trichloromethanethiol	16	(10	(16	(16	(16	(16
			(16	(16	(16	(16	(16 (10
B11	Trichloropropene	10	(18	(16	(16	(16	•
B12	1,2,3-Trichloropropene	16	(16	(16	(16	(16	(10
B15	Diethylarsine	16	(10	(18 (588	(16 (566	(16 (500	<16 <588
C71	Formaldehyde	580	(588				•
C#4	Methyl Methacrylate	16	(16	<10	(16	(16	(10
HØ6	Ethyl Mathacrylate	16	(16	(16	<16	(16	(10
819	Acetonitrile	3 ppm		(3 ppm	<3 ppe		(3 ppm
HØS	Ethylene Oxide	3 ppm	(3 ppm	<3 ppm	(3 ppm	(3 ppm	(3 ppm
733	Semi-Volatile Organic Compound						
B26	Aniline	< 16	< 18	< 16	< 16	< 16	< 10
B36	Benz (A) Anthracene	< 16	< 10	< 16	< 10	< 16	< 10
B33	Benzidine	< 16	< 16	< 10	< 16	< 16	< 16
B34	Benzo (B) Fluoranthene	< 10	< 10	< 10	< 16	< 16	< 10
B38	Bis (2-Chloroethoxy) Methane	< 10	< 10	< 16	< 10	< 16	< 10
839	Bis (2-Chloroethy!) Ether	< 10	< 10	< 10	< 16	< 10	< 18
848	Bis(2-Ethylhexyl)Phthalate	< 10	< 18	< 10	< 10	< 10	< 16
841	4-Brosophenyl Phenyl Ether	< 10	< 10	< 10	< 10	< 18	< 10
842	Butyl Ber.zyl Phthalate	< 10	< 10	< 10	< 10	< 10	(10 .
845	P-Chloroaniline	< 16	< 16	< 10	< 16	< 10	< 10
B46	P-Chloro M-Cresol	< 10	< 10	< 10	< 16	< 10	< 10
948	2-Chloronaphthalene	< 16	< 16	< 16	< 16	< 10	< 10
849	2-Chlorophenol	< 10	< 10	< 10	< 16	< 10	< 10
850	Chrysene	< 10	< 16	< 10	< 10	(16	< 10

TABLE B.7. (contd)

Analytical Code	Compound	Detection Limit (ppb)	88316 RM 8.9 Spring	88322 RM 14.5 Spring	88335 RM 28.1 Spring	88346 RM 42.1 Spring	88348 RM 42.3 Spring
733	Semi-Volatile Organic Compound	ds					
855	Dibenz (A, H) Anthracene	< 16	< 16	< 15	< 10	< 10	< 15
B68	Di-M-Buty! Phthalate	< 10	< 10	< 10	< 10	< 10	(15
B61	1,2-Dichlorobenzene	< 10	< 10	< 16	< 15	< 10	(15
862	1,3-Dichlorobenzene	< 10	< 10	< 10	< 10	< 10	< 10
B63	1,4-Dichlorobenzene	< 10	< 18	< 16	< 10	< 16	(15
B64	3,3'-Dichlorobenzidine	< 20	< 2€	< 28	< 20	< 25	< 20
B65	2,4-Dichlorophenol	< 10	< 10	< 10	< 18	< 10	< 10
B67	Diethyl Phthalate	< 16	< 16	< 18	< 10	< 10	(16
875	2,4-Disethylphenol	< 10	< 10	< 16	< 10	< 10 ·	(19
876	Disethyl Phthalate	< 16	< 15	< 10	< 10	(10	< 10
876	4,8-Dinitro-O-Cresol	< 10	< 10	< 16	< 10	< 10	< 10
B86	2,4-Dinitrotoluene	< 10	< 16	< 10	< 10	< 16	< 10
881	2,6-Dinitrotoluene	< 16	< 16	< 16	< 10	< 16	< 10
B82	Di-N-Octyl Phthalate	< 10	< 18	< 16	< 10	< 10	< 10
B84	1,2-Dipheny lhydrazine	< 10	< 16	< 16	< 10	< 10	< 10
888	Fluoranthene	< 16	< 16	< 16	< 15	< 16	< 10
889	Hexach lorobenzene	< 10	< 10	< 16	< 10	< 10	< 16
B96	Hexach lorobuted i ene	< 10	< 10	< 16	< 16	< 10	< 10
891	Hexachlorocyclopentadiene	< 10	< 16	< 16	< 10	< 10	< 16
892	Hexach loroethane	< 10	< 10	< 10	< 16	< 10	< 10
893	Indeno (1, 2, 3-CD) Pyrene	< 10	< 10	< 16	< 10	< 10	< 10
C54	Hexach lorophene	< 10	< 16	< 10	< 10	< 16	< 10
C55	Naphtha I ene	< 10	< 16	< 10	< 16	< 16	< 10
C12	Nitrobenzene	< 10	< 10	< 10	< 16	< 10	< 16
C17	N-Nitrosodimethylamin	< 10	< 10	< 16	(16	(10	< 10
C57	Phono I	< 16	< 16	< 16	< 16	< 10	< 15
C43	1,2,4-Trichlorobenzene	< 10	< 16	< 10	< 16	< 16	< 15
879	2,4-Dinitrophenol	< 5€	< 50	< 5€	< 50	< 5€	< 5€
C13	4-Nitrophenol	< 5€	< 50	< 5€	< 50	< 50	< 5€
C11	Para-Nitrosniline	< 5€	< 5€	< 5€	< 58	< 58	₹ 50
C28	Pentach loropheno l	< 50	< 58	< 5€	< 58	< 5€	< 5€
C44	2,4,5-Trichlorophenol	< 50	< 50	< 50	< 50	< 5€	< 50
C45	2,4,5-Trichlorophenol	< 16	< 10	< 16	< 16	< 10	< 10
C56	1,2,3-Trichlorobenzene	< 16	< 16	< 10	< 10	< 10	< 10
CS8	1,3,5-Trichlorobenzene	< 10	(15	< 18	< 15	< 10	(10
C37	1,2,4,5-Tetrachiorobenzene	< 10	< 16	< 10	< 10	< 10	< 10
C59	1,2,3,4-Tetrachiorobenzene	< 10	< 10	(10	(10	< 15	(16
CSS	1,2,3,5-Tetrachlorobenzene	(10	(16	< 16	< 10	< 16	< 10
C25	Pentach I orobenzene	< 16	< 18	< 16	< 10	< 10	< 10
851	Cresoia	(16	< 10	< 10	< 10	< 10	< 18
885	N-Nitrosodinpropylamine	< 10	< 10	< 18	< 18	< 18	< 18

TABLE B.7. (contd)

Ann luk!		Detection	88351	88353 Slide Seep	88355 Ringold	88357 Priest Rapid	88359 Richland
Analytica Code	Cospound	(ppb)	Byers Lndg Irr Rtn	Sirge Seeb	Irr Rtn	River	Pumphouse
	1200			********			
733	Semi-Volatile Organic Compound	ds					
B55	Dibenz (A, H) Anthracene	< 10	< 10	< 16	< 10	< 10	< 10
B66	Di-N-Butyl Phthalate	< 16	< 16	< 10	< 10	< 10	< 10
B61	1,2-Dichlorobenzene	< 16	< 16	< 10	< 10	< 16	< 16
B62	1,3-Dichlorobenzene	< 10	< 16	< 16	< 16	< 16	< 10
963	1,4-Dichlorobenzene	< 15	< 16	< 10	< 16	< 16	< 18
B64	3,3'-Dichlorobenzidine	< 26	< 26	< 26	< 26	< 26	< 20
B65	2,4-Dichlorophenol	< 16	< 10	< 16	< 16	< 10	< 16
B67	Diethyl Phthalate	< 16	< 16	< 16	< 16	< 16	< 16
875	2,4-Disethy Iphenol	< 15	< 16	< 16	< 10	< 10	< 16
B76	Disethyl Phthalate	< 10	< 10	< 16	< 18	₹ 16	< 16
878	4,6-Dinitro-O-Cresol	< 10	~ 16	< 16	< 10	< 10	< 10
B84	2,4-Dinitrotoluene	< 15	< 10	< 10	< 16	< 10	< 10
B81	2,5-Dinitrotoluene	< 10	< 10	< 15	< 10	< 16	< 16
B82	Di-N-Octyl Phthalate	< 16	< 16	< 16	< 16	< 15	< 15
B84	1,2-Diphenylhydrazine	< 16	< 16	< 16	< 16	< 15	< 16
B88	Fluoranthene	< 10	< 16	< 16	< 16	< 16	< 10
B89	Hexach lorobenzene	< 16	< 16	< 16	< 16	< 16	(-10
B96	Hexach lorobutadiene	< 10	< 16	< 16	< 10	< 16	< 16
891	Hexach lorocyc lopentad i ene	< 16	< 16	< 16	< 15	< 16	< 16
B92	Hexach loroethane	< 16	< 16	< 16	< 18	< 16	< 10
B93	Indeno(1,2,3-CD)Pyrene	< 10	< 16	< 18	< 16	< 16	< 16
C54	Hexach lorophene	< 10	< 18	< 16	< 16	< 16	< 16
CSS	Naphtha lene	< 16	< 16	< 16	< 16	· ··· (- 16	< 10
C12	Nitrobenzene	< 10	< 18	< 16	< 16	< 16	< 16
C17	M-Mitrosodimethylamin	< 10	< 10	< 16	< 16	< 10	< 16
C57	Pheno i	< 16	< 16	< 10	< 16	< 16	< 18
· C43	1,2,4-Trichlorobenzene	< 10	< 10	< 16	< 10	< 16	< 10
B79	2,4-Dinitrophenol	< 50	< 50 ℃	< 5€	< 5€	< 58 .	< 5€
C13	4-Nitrophenol	< 58	< 58	< 58	< 5€	< 56	< 5€
C11	Para-Nitroaniline	< 58	< 5€	< 50	< 50	< €#	< 5€
C28	Pentach loropheno l	< 58	< 50	< 56	< 58	< 58	< 5€
C44	2,4,5-Trichlorophenol	5.6	< 58	< 5€	< 58	< 56	< 5€
C45	2,4,6-Trichlorophenol	< 10	< 10	< 15	< 15	< 15	< 10
C56	1,2,3-Trichlorobenzene	< 10	< 15	< 10	< 16	< 10	< 10
C58	1,3,5-Trichlorobenzene	< 10	< 16	< 16	< 16	< 16	< 10
C37	1,2,4,5-Tetrachlorobenzene	< 16	< 16	< 16	< 16	< 10	< 16
C59	1,2,3,4-Tetrachlorobenzene	< 16	< 16	< 16	< 10	< 16	< 16
C66	1,2,3,5-Tetrachlorobenzene	< 10 .	< 15	< 16	< 16	< 10	< 16
C26	Pentach lorobenzene	< 10	< 10	< 10	< 15	< 15	< 10
B51	Cresois	< 10	< 16	< 15	< 10	< 10	< 10
885	N-Nitrosodinpropylamine	< 15	< 10	< 10	< 10	< 10	< 16

TABLE B.7. (contd)

Analytical		Detection	88316 RM 8.9	88322 RM 14.5	88335 RM 28.1	88346 RM 42.1	88348 RM 42.3
Code	Compound	(ppb)	Spring	Spring	Spring	Spring	Spring
733	Semi-Volatile Organic Compounds						
C49	Benzo (A) Pyrene	< 16	(10	(15	(10	(10	(10
CS1	Bis(2-Chloroisopropyl) Ether	(10	(19	(10	(10	(16	< 10
I21	Tributy phosphate	(10	< 10	(10	(16	(16	(10
B26	Acetophenone	(10	(18	(16	(16	(16	< 10
B21	Varfarin	< 10	< 10	(16	(10	(16	< 19
B22	2-Acetylaminofluorene	< 10	< 10	< 16	(16	< 10	< 10
B23	4-An inob i pheny l	< 10	< 10	(10	< 10	< 10	< 10
B24	5-(Aminomethyl)-3-Isoxazolol	< 10	< 10	(10	< 10	< 10	< 10
825	Anitrole	< 10	< 10	< 16	< 16	< 16	< 10
827	Aranita	< 10	< 15	< 16	< 15	< 10	< 10
B28	Auranine	< 10	< 16	(10	< 10	< 10	< 16
829	Benz(C)Acridine	< 16	< 10	< 16	< 10	< 19	< 15
B31	Benzene, Dichloromethyl	< 15	< 10	(16	(15	(10	< 10
B32	Benzesethoil	< 15	< 10	< 18	(10	< 10	(16
B36	P-Benzoquinone	(15	< 15	< 10	(16	(10	(10
837	Benzyl Chloride	(16	< 16	(15	(19	(18	(16
B43	2-Sec-Buty1-4,6-Dinitropheno	1	< 10	< 16	(10	< 19	< 10
B44	Chlorosikyl Ethers	< 15	(10	< 10	< 10	< 10	< 16
B47	1-Chioro-2,3-Epoxypropane	< 10	< 10	< 16	< 10	< 10	< 10
B52	2-Cyclohexyl-	< 10	(16	< 19	< 16	< 16	< 10
	4.6-Dinitrophenol						
B63	Dibenz (A, H) Acridine	< 15	< 16	< 16	< 10	< 15	< 10
R54	Dibenz (A, J) Acridine	< 15	< 15	< 16	< 16	< 15	< 10
B56	7H-Dibenzo (C, G) Carbazole	< 10	(16	< 16	< 16	< 10	< 10
B57	Dibenzo (A, E) Pyrene	< 10	< 10	< 10	< 15	< 16	< 10
858	Dibenzo (A, H) Pyrene	< 10	< 10	< 10	< 10	< 15	< 18
859	Dibenzo (A, I) Pyrene	< 15	< 15	< 16	< 10	< 10	< 18
B66	2,8-Dichlorophenol	< 10	< 15	< 16	< 10	< 16	< 10
968	Dihydrosafrole	< 10	< 10	< 10	< 10	< 10	< 15
B69	3,3'-Dimethoxybenzidine	< 10	< 15	< 10	< 18	< 10	< 10
876	P-Disethy Iss i noazobenzene	< 10	< 16	< 10	< 19	< 15	< 16
871	7,12-Disethylbenz(A) Anthracene	< 18	< 10	(10	< 16	(16	< 10
872	3,3'-Dimethy!benzidine	< 15	< 10	< 10	< 16	< 10	< 10
873	Thiofanox	< 10	< 15	< 10	< 15	< 16	< 10
874	Alpha, Alpha-Disethylphene- thylamine	< 10	< 16	< 16	(10	(10	(10
877	Dinitrobenzene	(16	< 15	< 10	< 16	< 10	< 10
883	Dipheny (seine	< 16	< 15	< 10	< 16	< 15	< 10
B86	Ethyleneisine	< 10	< 15	'< 10	< 10	< 10	< 10
B87	Ethyl Methanesulfonate	< 10	< 10	< 15	< 16	< 16	< 10
B94	Isosafrole	< 15	< 10	< 10	< 10	< 10	< 10

TABLE B.7. (contd)

Analytical Code	Compound	Detection Limit (ppb)	88351 Byers Lndg Irr Rtn	88353 Slide Seep	88355 Ringold Irr Rtn	88357 Priest Rapid River	88359 Richland Pumphouse
733	Semi-Volatile Organic Compounds						
C49	Benzo (A) Pyrene	< 16	< 10	< 15	< 16	< 10	< 15
CS1	Bis(2-Chloroisopropyl) Ether	< 16	< 16	< 16	< 16	< 10	< 18
I21	Tributy i phosphate	< 10	< 10	< 10	< 16	< 10	< 10
820	Acetophenone	< 16	< 16	< 15	< 16	< 16	< 10
B21	Warfarin	< 10	< 16	< 10	< 16	< 15	< 10
B22	2-Acety laminof luorene	< 18	< 18	< 10	< 10	< 18	< 10
823	4-Aminobiphenyl	< 10	< 10	< 16	< 10	< 10	< 10
824	5-(Aminomethyl)-3-Isoxazolol	< 10	< 10	< 10	< 10	< 16	< 15
B25	Amitrole	< 10	< 10	< 16	< 10	< 10	< 10
B27	Aranite	< 16	< 10	< 16	< 15	< 10	< 10
B28	Auranine	< 16	< 16	< 10 .	< 10	< 15	< 16
B29	Benz (C) Acridine	< 16	< 16	< 16	< 16	< 10	< 16
831	Benzene, Dichloromethyl	< 10	< 16	< 16	< 10	< 10	< 15
B32	Benzemetho i I	< 15	< 10	< 16	< 16	< 10	< 10
836	P-Benzoqu i none	< 16	< 16	< 10	< 18	< 15	< 16
B37	Benzyl Chloride	< 18	< 16	< 10	< 10	< 16	< 16
B43	2-Sec-Buty I-4,6-Dinitropheno	1 (10	< 10	< 16	< 10	< 10	< 10
B44	Chlorosikyl Ethers	< 15	< 16	< 10	< 16	< 10	< 15
B47	1-Chloro-2,3-Epoxypropane	< 15	< 10	< 15	< 15	< 15	₹ 16
852	2-Cyclohexyl- 4,6-Dinitrophenol	< 16	(10	< 10	< 10	< 15	< 15
863	Dibenz (A, H) Acridine	< 16	< 16	< 10	< 10	< 10	< 10
854	Dibenz(A, J) Acridine	< 16	< 16	< 10	< 15	< 10	< 10
B56	7H-Dibenzo(C,G)Carbazole	< 10	< 18	< 10	< 16	< 16	< 10
B57	Dibenzo (A, E) Pyrene	< 10	< 10	< 10	< 10	< 16	< 10
B58	Dibenzo (A, H) Pyrene	< 10	< 16	< 16	< 16	< 10	< 10
B59	Dibenzo (A, I) Pyrene	< 16	< 16	< 10	< 10	< 10	< 15
966	2,6-Dichtorophenol	< 10	< 10	< 10	< 10	< 10	< 19
868	Dihydrossfrole	< 15	< 16	< 10	< 10	< 15	< 10
869	3,3'-Dimethoxybenzidine	< 15	< 16	< 10	< 16	< 10	< 10
B7#	P-Disethy las i no azobenzene	< 10	< 10	< 10	< 15	< 16	< 10
871	7, 12-Disethylbenz (A) Anthracene	< 16	(18	< 10	< 10	< 16	(10
B72	3,3'-Dimethylbenzidine	< 10	< 10	< 10	< 10	< 10	< 10
873	Iniofanox	< 10	< 10	< 15	< 15	< 10	< 10
B74	Alpha, Alpha-Disethylphene- thylamine	< 16	< 18	(16	(10	< 10	< 10
877	Dinitrobenzene	< 10	< 10	< 10	< 10	< 10	< 10
883	Diphenylanine	< 15	< 16	< 10	< 10	< 15	< 10
886	Ethyleneisine	< 10	< 16	< 15	< 10	< 15	< 16
B87	Ethyl Methanesulfonate	< 15	< 16	< 15	< 10	< 10	< 16
B94	Isosafrole	< 10	< 10	< 15	< 18	< 10	< 10

TABLE B.7. (contd)

		Detection	88316	88322	88335	88346	88348
Analytical		Linit	RM 8.9	RM 14.5	RM 28.1	RM 42.1	RM 42.3
Code	Compound	(ppb)	Spring	Spring	Spring	Spring	Spring
733	Semi-Volatile Organic Compound	•					
B95	Melononitrile	< 10	< 10	< 10	< 16	< 10	< 18
B96	Malphalan .	< 10	< 10	< 10	< 10	< 10	< 10
897	Mathapyri lene	< 15	< 15	< 10	< 16	< 10	< 19
898	Matho lony I	< 10	< 16	< 10	< 16	< 18	< 16
B99	2-Methylaziridine	< 10	< 10	< 10	< 16	< 10	< 10
CS1	3-Methylcholanthrene	< 10	< 10	< 10	< 10	< 10	< 10
C#2	4,4'-Methy lenebis (2-Chloro- aniline)	< 10	< 10	< 10	< 16	< 10	< 16
CS3	2-Methyllactonitrile	< 15	< 10	< 15	< 16	< 16	< 10
COS	Methy! Methanesulfonate	< 15	< 10 ·	< 16	< 16	< 16	< 10
C86	2-Methy I-2- (Methy I thio) Prop- ional dehyde-5- (Methy I- carbony I) 0xise	< 16	〈 10	< 16	< 16	< 10	₹ 10
C07	Methy Ith iourscil	< 10	< 16	< 15	< 16	< 16	< 10
CS8	1,4-Naphthoquinone	< 15	< 16	< 19	< 10	< 10	< 15
CS9	1-Napthy lasine	< 10	< 16	< 15	< 16	< 10	< 10
C16	2-Hapthy lasine	< 15	< 15	< 16	< 16	< 16	< 16
C14	N-Nitrosodi-N-Butylamine	< 15	< 16	< 16	< 16	< 10	< 16
C15	N-Nitrosodiethanolazine	< 10	< 10	< 10	< 16	< 16	< 10
C16	N-Nitrosodiethy lamine	< 16	< 16	< 15	< 16	< 10	< 16
C18	M-Nitrosomethy lethy lamine	< 15	< 16	< 15	< 16	< 16	< 16
C19	N-Nitroso-N-Methylurethane	< 15	< 15	< 15	< 15	(16	< 16
C26	N-Nitrosomethy lviny lamine	< 19	< 15	< 18	< 15	< 10	< 16
C21	N-Nitrosomorpholine	< 10	< 10	< 18	< 10	< 10	< 16
C22	N-Nitrosonornicotine	< 15	< 10	< 16	< 15	< 15	< 16
C23	N-Nitrosonopiperidine	< 15	< 15	< 15	< 10	< 15	< 16
C24	Mitrosopyrrolidine	< 19	< 15	< 15	< 10	< 15	< 16
C25	5-Mitro-O-Toluidine	< 18	< 10	< 16	< 10	< 16	< 16
C27	Pentach loron i trobenzene	< 18	< 10	< 16	< 16	< 10	< 16
C29	Phenacetin	< 15	< 10	(18	< 16	< 10	< 10
C36	Pheny lened iss ine	< 15	< 16	< 10	< 10	< 15	< 15
C31	Phthalic Acid Esters	< 10	< 16	< 10	< 10	< 16	< 10
C32	2-Picoline	< 15	< 16	< 15	< 15	< 16	< 16
C33	Pronamide	< 16	< 16	< 15	< 16	< 16	< 10
C34	Reserpine	< 16	< 10	< 10	< 16	< 16	< 10
C35	Rescorzinol	(15	< 16	< 16	< 15	< 16	< 15
C36	Safrol	< 15	< 10	< 10	< 15	< 16	< 18
C39	2,3,4,6-Tetrachlorophenol	< 15	< 15	< 10	(16	< 16	< 15
C48	Thiurae	< 16	< 16	< 10	< 10	< 10	< 16
C41	Toluenedianine	< 16	< 16	< 15	< 10	< 18	(10
C42	O-Toluidine Hydrochloride	< 16	< 16	< 10	< 16	< 10	< 16

TABLE B.7. (contd)

Analytical Code	Compound	Detection Limit (ppb)	Irr Rtn	88353 Slide Seep	88355 Ringold Irr Rtn	88357 Priest Rapid River	Pumphouse
733	Semi-Volatile Organic Compound	6		***************************************			
B95	Malononitrile	(10	< 10	< 18	< 15	< 10	< 10
B96	Malphalan	< 16	< 10	< 10	< 10	< 10	< 10
B97	Mathapyrilene	< 1€	< 10	< 10	< 10	< 10	< 10
898	Matholonyl	< 15	< 10	< 10	< 10	< 16	< 16
B99	2-Methy laziridine	< 18	< 10	< 10	< 18 ·	< 10	< 10
CØ1	3-Methylcholanthrene	< 10	< 15	< 10	< 15	< 10	< 10
CS2	4,4'-Methylenebis(2-Chloro- aniline)	(10	< 10	< 10	(10	< 10	< 19
C#3	2-Methyllactonitrile	< 16	< 16	< 10	< 15	< 10	< 10
CØ5	Methy! Methanesulfonate	< 10	< 10	(10	< 18	< 10	< 18
C\$6	2-Methyl-2-(Methylthio)Prop- ionaldehyde-6-(Methyl- carbonyl)Oxime	< 10	(10	< 10	< 1€	< 1€	< 18
C67	Methy Ithiourscil	< 10	< 10	< 16	< 10	< 1€	< 10
C#8	1,4-Naphthoguinone	< 10	< 10	< 16	< 18	< 10	(18
C#9	1-Napthy lamine	< 10	< 10	< 16	< 10	< 10	< 18 ·
C18	2-Napthy lanine	< 10	< 10	< 18	< 10	< 10	< 18
C14	N-Nitrosodi-N-Butylanine	< 16	< 10	< 15	< 18	< 10	< 10
C15	N-Nitrosodiethanolazine	(15	< 10	< 16	< 10	< 10	< 18
C16	N-Nitrosodiethylamine	< 10	(16	< 10	< 10	< 10	< 18
C18	N-Nitrosomethy lathy lamine	< 10	< 10	< 10	< 18	< 10	< 10
C19	N-Nitroso-N-Methylurethane	< 10	< 10	< 10	< 10	< 10	< 10
C29	N-Nitrosomethylvinylamine	< 10	< 10	< 10	< 10	< 16	< 18
C21	N-Nitrosomorpholine	< 10	< 18	< 16	< 10	< 10	(15
C22	N-Nitrosonornicotine	< 10	< 10	< 10	< 10	< 10	< 16
C23	N-Nitrosonopiperidine	< 16	< 10	< 16	< 10	< 10	< 10
C24	Nitrosopyrrolidine	< 16	< 10	< 10	< 10	< 10	< 16
C25	5-Nitro-O-Toluidine	< 10	< 16	< 10	< 10	< 10	< 18
C27	Pentachloronitrobenzene	< 10	< 18	< 10	< 16	< 16	< 10
C29	Phenacetin	< 10	< 1€	< 10	< 16	< 16	< 16
C36	Phenylenedismine	< 10	< 10	< 10	< 10	< 10	< 15
.C31	Phthalic Acid Esters	< 16	< 16	< 10	< 16	< 16	< 10
C32	2-Picoline	< 10	< 10	< 16	< 16	< 10	< 10
C33	Pronanide	< 16	< 10	< 10	< 10	< 16	< 10
C34	Reserpine	< 10	< 10	< 10	< 1s	< 18	< 10
C35	Rescorzinol	< 10	< 10	< 10	< 16	< 19	< 18
C36	Safroi	< 10	< 10	< 10	< 16	< 16	< 19
C39	2,3,4,6-Tetrachlorophenol	< 10	< 10	< 16	< 16	(10	< 19
C48	Thiurae	< 10	< 10	< 10	< 16	< 18	< 16
C41	Toluenedismine	< 16	< 16	< 15	< 16	< 18	< 10
C42	O-Toluidine Hydrochloride	< 19	< 10	< 19	< 10	< 15	< 10

TABLE B.7. (contd)

Analytical		Detection Limit	88316 RM 8.9	88322 RM 14.5 Spring	88335 RM 28.1 Spring	88346 RM 42.1 Spring	18348 Rd 42.3 Spring
Code	Compound	(ppb)	Spring	Spring	Spring	spring	
733	Semi-Volatile Organic Compour	nds					
C48	0,0,0-Triethyl Phosphoro- thiosts	< 10	< 10	< 16	< 10	(10	(10
C47	Sym-Trinitrobenzene	< 18	< 10	< 10	< 16	< 10	< 10
C48	Tris(2,3-Dibrosopropy1)- Phosphate	(10	(18	< 10	(10	< 10	(10
CSO	Chloronaphazine	< 10	< 15	< 10	< 10	< 15	< 10
C52	Hexach loropropene	< 10	< 10	< 10	< 16	< 16	< 10
C79	Kerosene	< 10	< 10	< 10	< 10	< 10	< 16
C92	Maleic Hydrazide	<580	<500	(586	(566	(586	<566
C93	Nicotinic Acid	(188	<188	<186	<166	<166	<100
C91	Strychnine	< 50	< 50	< 50	< 5€	< 5€	< 50
B35	Benzo(J)Fluoranthene	< 10	< 16	< 18	< 10	< 10	< 10
734	Phosphorous Pesticides						
C81	Tetraethy pyrophosphate	2	<2	<2	<2	(2	<2
C63	Carbophenothion	2	<2	<2	(2	<2	<2
C84	Disulfoton	2	<2	<2	(2	(2	<2
C85	Disethoate	2	<2	<2	(2	(2	⟨2
C66	Methylparathion	2	(2	<2	<2	(2	<2
C67	Ethyl Parathion	2	(2	(2	(2	(2	⟨2
736	Direct Aqueous Injection Ana	lysis					
C53	Hydrazine	3000	(3000	<3966	<3966	<3886	<3666
C98	Para I dehyde	2006	<2 866	(2606	<2000	<20 00	<2666
C94	Acrylanide	19960	(18686	<1 6066	<18866	(15556	<18666
C95	Allyi Alcohoi	2500	(2566	(2500	⟨25€€	(2566	<2586
C97	Chloroacetaldehyde	16000	(16886	<16 666	(16666	<16888	<16959
C98	3-Chloropropionitrile	4966	<4000	<4 886	<4586	<4 988	<4866
HØ3	Ethyl Carbanate	5000	<5000	<5000	(5898	<5888	<5000
HØ4	Ethyl Cyanide	2866	<2686	<2866	(2555	<2888	(2866
H#9	Isobuty Alcohol	1996	<1666	<1866	<1888	<1866	<1000
H11	N-Propylamine	15006	(16666	(10000	<100 00	<10000	<1 6666
H12	2-Propyn-1-0L	8000	(8666	(8996	(8986	<8 956	(8866
737	Enhanced Herbicides						
H13	2,4-0	2	<2	(2	<2	<2	<2
H14	2(2,4,5-T)P	2	<2	<2	<2	(2	<2
H15	(2,4,5-T)	2	<2	<2	(2	<2	<2

TABLE B.7. (contd)

Analytica Code	Coapound	Detection Limit (ppb)	88351 Byers Lndg Irr Rtn	88353 Slide Seep	88355 Ringold Irr Rtn	88357 Priest Rapid River	88359 Richland Pumphouse
733	Semi-Volatile Organic Compour	nda	***********		**********		
C48	0,0,0-Triethy! Phosphoro- thioste	⟨ 10	(16	< 15	(16	(16	< 15
C47	Sym-Trinitrobenzene	< 16	< 15	< 15	< 10	< 18	< 16
C48	Tris(2,3-Dibrosopropyl)- Phosphate	< 16	< 10	< 15	< 10	< 15	< 16
C50	Chloronaphazine	10	< 10	< 10	< 16	< 18	< 10
C52	Hexach l'oropropene	< 16	< 10	< 16	< 10	< 10	< 10
C79	Kerosene	< 10	< 15	< 16	< 10	< 15	< 15
C92	Maleic Hydrazide	<500	<586	(598	<500	(588	<500
C93	Nicotinic Acid	<100	<100	<186	<186	<100	<100
C91	Strychnine	< 5€	< 5€	< 5€	< 5€	< 5€	< 5€
B35	Benzo(J)Fluoranthene	< 10	< 15	< 1€	< 10	< 18	< 10
734	Phosphorous Pesticides						
C61	Tetraethy pyrophosphate	2	<2	<2	<2	<2	<2
C63	Carbophenothion	2	<2	<2	<2	<2	<2
C84	Disulfoton	2	<2	<2	(2	<2	<2
CSS	Disethoste	2	<2	<2	(2	<2	<2
C88	Methy parathion	2	<2	<2	<2	<2	(2
C87	Ethyl Parathion	2	⟨2	<2	(2	<2	(2
736	Direct Aqueous Injection Anal	ysis					
C53	Hydrazine	3006	<306€	(3666	<3666	<3866	<3000
C96	Paraldehyde	2000	(2000	<2866	<2000	<2000	<2000
C94	Acrylamide	10000	<10000	<18600	<18866	<10000	<10000
C95	Ally! Alcohol	2500	⟨256€	(2500	⟨25#€	(2566	<2569
C97	Chloroscetaldehyde	18000	<16606	(16666	<16666	<16000	<16009
C98	3-Chloropropionitrile	4900	<4000	<486E	<4666	<4000	<4886
HØ3	Ethyl Carbanate	5000	<5666	<5000	<5986	<5000	<5000
H64	Ethyl Cyanide	2000	<2000	<2506	<2000	<2000	<2000
HØ9	Isobuty! Alcoho!	1066	<1000	<1900	<1000	<1986	<1000
H11	N-Propylamine	19666	<18666	<10000	<10000	<10000	<10000
H12	2-Propyn-1-OL	8000	<8666	(8866	<8666	(8696	<8000
737	Enhanced Herbicides						
H13	2,4-D	2	<2	<2	<2	<2	<2
H14	2(2,4,5-T)P	2	⟨2	<2	<2	<2	<2
H15	(2,4,5-T)	, 2	<2	<2 ⋅	<2	<2	<2

TABLE B.7. (contd)

		Detection	88316	88322	88335	88346	88348
Analytica	l .	Limit	RM 8.9	RM 14.5	RM 28.1	RM 42.1	RM 42.3
Code	Compound	(ppb)	Spring	Spring	Spring	Spring	Spring
739	PCB Analysis						
A54	Arochior 1816	4	<1	(1	(1		(1
A55	Arochior 1221	1	<1	<1	<1		<1
A56	Arochier 1232	1	<1	<1	<1		<1
A57	Arochlor 1242	1	<1	<1	<1		<1
A58	Arochlor 1248	1	<1	<1	<1		<1
A59	Arochior 1254	1	<1	<1	<1		<1
ASS	Arochlor 1266	. 1	<1	<1	<1		<1
A2S	Arsenic	5	(\$	< 5	<5	<5	<5
A21	Mercury	6.1	<6.1	<0.1	⟨€.1	⟨€.1	< 6.1
A22	Selenium	5	< 5	< 5	<5	(5	<5
A23	Thallium	8	(5	(5	<5	<5	(5
A51	Lead	8	(5	(5	(5	(5	(5
C59	TOC (Total Organic Carbon)		486	1216	433	656	782
C76	Cyanide	16	<15	<16	16.5	(10	<16
C77	Perchlorate	1000	(1666	<1998	(1886	(1886	<1966
C78	Sulfide	1000	(1886	<1000	(1886	(1886	(1986
C86	Assonius Ion	56		(58	(56	(56	⟨5€
C81	Ethylene Glycol	16 ppm	(16 ppm	(16 ppm	(16 ppe	(18 ppm	(16 ppm
C87	Citrus Red	1000	(1886	<1866	<1866	<1866	(1666
H16	TC (Total Carbon)	2000	14558	18213	25468	14298	15718
H42	TOX (Total Organic Halogen)	16	<16		<18	36.2	24.9

TABLE B.7. (contd)

Analytical		Detection Limit	88351 Byers Lndg	88353 Slide Seep	88355 Ringold	88357 Priest Rapid	88359 Richland
Code	Compound	(ppb)	Irr Rtn		Irr Rtn	River	Pumphouse
739	PCB Analysis						
A54	Arochior 1515	4				<1	<1
A55	Arochior 1221	1				<1	<1
A56	Arochlor 1232	1				<1	<1
A57	Arochlor 1242	1				<1	<1
A58	Arochlor 1248	1				<1	<1
A59	Arochior 1254	1				(1	<1
ASE	Arochior 1260	1				<1	<1
A25	Arsenic	5	6	(5	6	(5	⟨5
A21	Mercury	0.1	⟨∅.1	<6.1		<5.1	⟨∅.1
A22	Selenium	5	<5	16	5	(5	< 5
A23	Thallium	5	<5	<5	<5	₹5	(5
A51	Lord	5	<5	<5	<5	(5	(5
C69	TOC (Total Organic Carbon)		2666	2900	2500	1281	1272
C78	Cyanide	10	<10	<10	<10	(16	<1€
C77	Perchlorate	1000	<1696	<1000	<1000	<1500	<1000
C78	Sulfide	1000	<1000	<1000	<1566	<1800	<1000
C86	Assonius Ion	50	<5€	<5€	⟨5#	<5€	(50
C81	Ethylene Glycol	16 ppm	<16 ppm	(16 ppm	<16 ppm	<10 pps	(16 ppm
C87	Citrus Red	1666	<1886	<1466	<1966	<1966	<1666
H16	TC (Total Carbon)	2886	46776	39928	47296	13326	14163
H42	TOX (Total Organic Halogen)	10	28	12	14	<10	<10

TABLE B.7. (contd)

		Detection	88385	88316	88319	88326	88322
Analytica		Limit	RM 3.3	RM 8.9	RM 11.6	RM 14.5	RM 14.5
Code	Compound	(ppb)	Spring	Spring	Spring	River	Spring
735	IC Report						
C72	Nitrate	586	6766	28536	2266	(586	<596
C73	Sulphate	5666	36966	35576	13866	16147	13167
C74	Fluoride	560	(500	<566	<566	(566	<566
C75	Chloride	5000	6786	1879	1956	850	1113
C76	Phosphate	1666	(1969	(1556	<1866	<1566	<1666
		Detection	88328	88336	88332	88333	88335
Analytica	ŧ	Linit	RM 27.75	RM 27.25	RM 27.5	RM 28.1	RM 28.1
Code	Compound	(ppb)	Spring	Spring	Spring	River	Spring
		(ppo)	opr mg	opt tilg			opi mg
735	IC Report						
C72	Nitrate	586	2297	6363	12713	31296	31646
C73	Sulphate	5000	14872	24767	33416	36546	38366
C74	Fluoride	500	<566	(586	<566	<500	<500
C75	Chloride	5000	1942	4999	6396	9246	9116
C76	Phosphate	1986	(1886	(1866	(1866	(1866	<1866
		Detection	88336	88338	88340	88341	88342
Analytica	1	Linit	RM 28.5	RM 2.5	RM 38.25	RM 38.8	RM 41.58
Code	Compound	(ppb)	Spring	Spring	Spring	Spring	Spring
705							*********
735 C72	IC Report Nitrate	566	28288	800	25437	13691	4884
C72	Sulphate	5866	38486	18986	28349	36836	28894
C74	Fluoride	566	(586	(588	₹556	(566	(588
C74	Chloride	5000	7658	866	15985	9588	7766
C78		1666		(1966	(1995	(1666	(1888
C/6	Phosphate	1999	<1000	(1000	(100)	(1000	(1000

TABLE B.7. (contd)

		Detection	88343	88344	88346	88348	88349	
Analytical		Lieit	RM 42.6	RM 42.1	RM 42.1	RM 42.3	RM 43.6	
Code	Compound	(ppb)	Spring	River	Spring	Spring	Spring	

735	IC Report							
C72	Nitrate	500	28458	2149	1697	9183	9166	
C73	Sulphate	5666	47346	17276	17423	16326	14851	
C74	Fluoride	566	(586	(566	<586	(586	(588	
C75	Chloride	5666	15426	6926	7586	13470	2573	
C76	Phosphate	1500	(1666	(1666	(1996	<1000	(1666	
		Datection	88351	86353	88355	88367	88359	
Analytical		Linit	Byers Lndg	Slide Seep	Ringold	Priest Rapid	Richland	
Code	Compound	(ppb)	Irr Rtn		Irr Rtn	River	Pumphouse	
735	IC Report							
C72	Nitrate	500	15631	5232	16795	<566	554	
C73	Sulphate	5888	56676	238666	63766	16336	16862	
C74	Fluoride	566	(586	<500	<588	(586	(500	
C75	Chloride	5466	13560	42370	15816	895	1619	
C76	Phosphate	1966	(1966	<1000	<1986	<1000	<1888	